



Designing global climate and atmospheric chemistry simulations for 1 and 10 km diameter asteroid impacts using the properties of ejecta from the K-Pg impact

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Abstract. About 66 million years ago, an asteroid about 10 km in diameter struck the Yucatan Peninsula creating the Chicxulub crater. The crater has been dated and found to be coincident with the Cretaceous–Paleogene (K-Pg) mass extinction event, one of six great mass extinctions in the last 600 million years. This event precipitated one of the largest episodes of rapid climate change in Earth's history, yet no modern three-dimensional climate calculations have simulated the event. Similarly, while there is an ongoing effort to detect asteroids that might hit Earth and to develop methods to stop them, there have been no modern calculations of the sizes of asteroids whose impacts on land would cause devastating effects on Earth. Here, we provide the information needed to initialize such calculations for the K-Pg impactor and for a 1 km diameter impactor.

There is considerable controversy about the details of the events that followed the Chicxulub impact. We proceed through the data record in the order of confidence that a climatically important material was present in the atmosphere. The climatic importance is roughly proportional to the optical depth of the material. Spherules with diameters of several hundred microns are found globally in an abundance that would have produced an atmospheric layer with an optical depth around 20, yet their large sizes would only allow them to stay airborne for a few days. They were likely important for triggering global wildfires. Soot, probably from global or near-global wildfires, is found globally in an abundance that would have produced an optical depth near 100, which would effectively prevent sunlight from reaching the

surface. Nanometer-sized iron particles are also present globally. Theory suggests these particles might be remnants of the vaporized asteroid and target that initially remained as vapor rather than condensing on the hundred-micron spherules when they entered the atmosphere. If present in the greatest abundance allowed by theory, their optical depth would have exceeded 1000. Clastics may be present globally, but only the quartz fraction can be quantified since shock features can identify it. However, it is very difficult to determine the total abundance of clastics. We reconcile previous widely disparate estimates and suggest the clastics may have had an optical depth near 100. Sulfur is predicted to originate about equally from the impactor and from the Yucatan surface materials. By mass, sulfur is less than 10 % of the observed mass of the spheres and estimated mass of nanoparticles. Since the sulfur probably reacted on the surfaces of the soot, nanoparticles, clastics, and spheres, it is likely a minor component of the climate forcing; however, detailed studies of the conversion of sulfur gases to particles are needed to determine if sulfuric acid aerosols dominated in late stages of the evolution of the atmospheric debris. Numerous gases, including CO₂, SO₂ (or SO₃), H₂O, CO, Cl, Br, and I, were likely injected into the upper atmosphere by the impact or the immediate effects of the impact such as fires across the planet. Their abundance might have increased relative to current ambient values by a significant fraction for CO₂, and by factors of 100 to 1000 for the other gases.

For the 1 km impactor, nanoparticles might have had an optical depth of 1.5 if the impact occurred on land. If the

impactor struck a densely forested region, soot from the forest fires might have had an optical depth of 0.1. Only S and I would be expected to be perturbed significantly relative to ambient gas-phase values. One kilometer asteroids impacting the ocean may inject seawater into the stratosphere as well as halogens that are dissolved in the seawater.

For each of the materials mentioned, we provide initial abundances and injection altitudes. For particles, we suggest initial size distributions and optical constants. We also suggest new observations that could be made to narrow the uncertainties about the particles and gases generated by large impacts.

1 Introduction and definitions

About 66 million years ago, an asteroid around 10 km in diameter hit the Earth near the present-day Yucatan village of Chicxulub and created an immense crater whose age coincides with the Cretaceous–Paleogene (K–Pg) global mass extinction (Alvarez et al., 1980; Schulte et al., 2010; Renne et al., 2013). There is an enormous literature concerning this event and its aftermath. Surprisingly, however, there are very few papers about the changes in climate and atmospheric chemistry caused by the debris from the impact while it was in the atmosphere, and no studies based on modern three-dimensional climate models. Nevertheless, this event was almost certainly one of the largest and most dramatic short-term perturbations to climate and atmospheric chemistry in Earth's history.

There is substantial evidence for many other impacts in Earth's history as large or larger than that at Chicxulub, mostly in the Precambrian (e.g., Johnson and Melosh, 2012a; Glass and Simonson, 2012). There is also a growing effort to find asteroids smaller than the one that hit Chicxulub, but whose impact might have significant global effects, and to develop techniques to stop any that could hit the Earth. For example, as of 17 November 2015, NASA's Near Earth Object Program identifies 13 392 objects whose orbits pass near Earth. Among these objects, 878 have a diameter of about 1 km or larger, and 1640 have been identified as "potentially hazardous asteroids", which are asteroids that pass the Earth within about 5% of Earth's distance from the sun, and are larger than about 150 m diameter.

There is evidence for such smaller impacts in recent geologic history from craters, osmium variations in sea cores (Paquay et al., 2008), and spherule layers (Johnson and Melosh, 2012a; Glass and Simonson, 2012). For instance, a multi-kilometer object formed the Siberian Popigai crater in the late Eocene and another multi-kilometer object formed the late Eocene Chesapeake Bay crater in the United States. Size estimates vary between techniques, but within a given technique the Popigai object is generally given a diameter half that of the Chicxulub object. Toon et al. (1997) point out

that the environmental effects of impacts scale with the impactor energy, or cube of the diameter, not diameter (or crater size). The Popigai object likely had about 12% of the energy of the Chicxulub object. Surprisingly, except for collisions in the ocean (Pierazzo et al., 2010), climate models have not been used to determine the destruction that might be caused by objects near 1 km in diameter, a suggested lower limit to the size of an impactor that might do significant worldwide damage (e.g., Toon et al., 1997).

Here, we describe the parameters that are needed to initialize three-dimensional climate and atmospheric chemistry models for the Chicxulub impact and for a 1 km diameter asteroid impact. Nearly every aspect of the K–Pg impact event is uncertain and controversial. We will address some of these uncertainties and controversies and make recommendations for the initial conditions that seem most appropriate for a climate model, based upon the geological evidence. We will also suggest the properties of the initial impact debris from a 1 km diameter asteroid.

There are numerous observed and predicted components of the Chicxulub impact debris. The distal debris layer, defined to be the debris that is more than 4000 km removed from the impact site, is thought to contain material that remained in the atmosphere long enough to be globally distributed. This distal layer, sometimes called the fireball layer or the magic layer, is typically only a few millimeters thick (Smit, 1999). As discussed below, the layer includes 200 μm sized spherules, 50 μm sized shocked quartz grains, 0.1 μm sized soot, and 20 nm sized iron-rich material.

We discuss each of the components of the distal layer in detail below. In brief, we find the following: the large spherules are not likely to be of importance to the climate because they would have been removed from the atmosphere in only a few days. However, they may have initiated global wildfires. The shocked quartz grains, one of the definitive pieces of evidence for an impact origin as opposed to volcanic origin of the debris layer, are likely only a small fraction of the clastic debris. It is difficult to identify the rest of the minerals produced by crushing because there is material in the layer that might have been produced long after the impact by erosion and chemical alteration of the large spheres or from the ambient environment. One major controversy surrounding the clastic material is the fraction that is submicron sized. Particles larger than a micron will not remain in the atmosphere very long and, therefore, are less likely to affect climate. Unfortunately, the submicrometer portion of the clastics in the distal layer, which might linger in the atmosphere for a year or more, has not been directly measured. Our estimate of the mass of submicron-sized clastics suggest that it could have had a very large optical depth that would be capable of modifying the climate significantly. Nevertheless, submicron clastics are only of modest climatic importance relative to the light-absorbing soot and possibly the iron-rich nanometer-scale debris. Submicron soot is observed in the global distal layer in such quantity that it would have had a

very great impact on the climate when it was suspended in the atmosphere. The major controversy surrounding the soot is whether it originated from forest fires or from hydrocarbons at the impact site. The origin of the soot, however, is of secondary importance with regard to its effect on climate. Since the soot layer overlaps the iridium layer in the distal debris, it had to have been created within a year or two of the impact, based on the removal time of small particles from the atmosphere (and ocean), and could not have been the result of fires long after the impact. The fireball layer is often colored red and contains abundant iron. Some of the iron has been identified as part of a 20 nm sized particle phase, possibly representing a portion of the recondensed vaporized impactor and target. However, relatively little work has been done on this material. Its abundance has not been measured, but theoretical work suggests its mass could have been comparable to that of the impactor. Therefore, the nanometer-sized particles could have been of great importance to the climate. Each of the materials just described is present in the distal layer, and their impacts on the atmosphere were likely additive.

There are several other possible components of the distal layer that have not been clearly identified and studied as part of the impact debris, which we discuss below. Water, carbon, sulfur, chlorine, bromine, and iodine were likely present in significant quantities in the atmosphere after the impact. The Chicxulub impact occurred in the sea with depths possibly ranging up to 1 km. The target sediments and the asteroid probably also contained significant amounts of water. Water is an important greenhouse gas, and could condense to form rain, which might have removed materials from the stratosphere. Carbon is present in seawater, in many asteroids, and in sediments. Injections as carbon dioxide or methane might have led to an increased greenhouse effect. Sulfur is widely distributed in the ambient environment and is water soluble. Therefore, it is difficult to identify extraterrestrial sulfur in the debris layer. However, the impact site contains a lot of sulfur, and asteroids also contain significant amounts of sulfur. Sulfur is noteworthy because it is known to produce atmospheric particulates in today's atmosphere that alter the climate. Chlorine, bromine, and iodine can destroy ozone, and their effectiveness as catalysts is enhanced by heterogeneous reactions on sulfuric acid aerosols.

In addition to the millimeter-thick distal layer, there is an intermediate region ranging from 2500 to 4000 km from the impact site with a debris layer that is several centimeters thick (Smit, 1999). This layer contains microtektites (molten rock deformed by passage through the air), shocked quartz, as well as clastics such as pulverized and shocked carbonates. Most of this layer originated from the target material in the Yucatan. It is of interest because, like the debris clouds from explosive volcanic eruptions, components of this material may have escaped from the region near the impact site to become part of the global debris layer.

Properties of each of these materials need to be known in order to model their effects on the climate and atmospheric

chemistry realistically. These properties include the altitude of injection, the size of the injected particles, the mass of injected particles or gases, the density of the particles, and the optical properties of the injected particles and gases. Our best estimates for these properties for the K-Pg impact are summarized in Table 1 for particles and Table 2 for gases, and discussed for each material in Sect. 2. Tables 3 and 4 provide an extrapolation of these properties for an impact of a 1 km sized object.

While the mass of the injected material is useful as an input parameter to a model, the optical depth of the particles is needed to quantify their impact on the atmospheric radiation field and, therefore, on the climate. Hence, optical depth is a useful quantity to compare the relative importance of the various materials to the climate. For a monodisperse particle size distribution, the optical depth is given by $\tau = \frac{Mq_{\text{ext}}}{4\rho r}$. Here, M is the mass of particles in a column of air (for example, g cm^{-2}), r is the radius of the particles, ρ is the density of the material composing the particles, and q_{ext} is the optical extinction efficiency at the wavelength of interest. The optical extinction efficiency is a function of the size of the particles relative to the wavelength of light of interest, and of the optical constants of the material. The optical extinction efficiency is computed accurately in climate models. However, a rough value of q_{ext} for particles larger than $1 \mu\text{m}$ is about 2 for visible wavelength light. We use this rough estimate for q_{ext} in Tables 1 and 3 to calculate an optical depth for purposes of qualitatively comparing the importance of the various types of injected particles. We assume in the heuristic calculations of optical depth in Tables 1 and 3 that the particles have a radius of $1 \mu\text{m}$ because smaller particles will quickly coagulate to a radius near $1 \mu\text{m}$ given the large masses of injected material. Particles smaller than $1 \mu\text{m}$ would lead to a larger optical depth than given in Tables 1 and 3.

Below, we define the properties that are needed to perform climate or atmospheric chemistry simulations for each material that might be important.

2 Particulate injections

2.1 Large spherules

2.1.1 Large spherules from the Chicxulub impact

The most evident component of the distal and regional debris layers is spherical particles, some of which are large enough to be seen with the naked eye. Due to their spherical shape it is assumed that they are part of the melt debris from the impact or the condensed vapor from the impact (Johnson and Melosh, 2012b, 2014). The particles are not thought to have melted on reentry into the atmosphere since debris launched above the atmosphere by the impact should not reach high enough velocities to melt when it reenters the atmosphere. According to Bohor and Glass (1995), there are

Table 1. K-Pg injection scenario for impactor mass $\sim 1.4 \times 10^{18}$ g; impact energy $\sim 2.8 \times 10^{23}$ J = 6.8×10^7 Mt for 20 km s^{-1} impact.

Property/ constituent	Type 2 spherules	Soot	Nanoparticles	Clastics, $< \mu\text{m}$	S
Material amount, g, column density (g cm^{-2})	2.3×10^{18} (0.44)	$1.5\text{--}5.6 \times 10^{16}$ to 1.1×10^{-2} ^c	$\sim 2 \times 10^{18\text{b}}$ (0.4)	$< 6 \times 10^{16}$ (0.01)	9×10^{16} ($5 \times 10^{-2} \text{ g cm}^{-2}$ as SO_4)
Global optical depth as 1 μm particles ^a	~ 20 (for 250 μm particles)	~ 100	~ 2000	~ 90	~ 450
Vertical distribution	70 km, Gaussian distribution with half width of 6.6 km ^d	Eq. (2)	Same as Type 2 spherules	Uniformly mixed vertically above tropopause	Same as Type 2 spherules
Optical properties	Not relevant	$n = 1.8$ $k = 0.67$	Hervig et al. (2009)	Orofino et al. (1998) limestone	Sulfuric acid
Initial particle size	250 μm diameter	Lognormal, $r_m = 0.11 \mu\text{m}$, $\sigma = 1.6$; monomers 30–60 nm	20 nm diameter	Lognormal, $r_m = 0.5 \mu\text{m}$, $\sigma = 1.65$	Gas
Material density, g cm^{-3}	2.7	1.8	2.7	2.7	1.8

^a Qualitative estimate for comparison purposes only. ^b This value is an upper limit. The lower limit is zero. ^c These values are for aciniform soot or elemental carbon in the stratosphere (see text). ^d The material may have quickly moved to below 50 km to maintain hydrostatic balance (see text).

two types of spherules, with differing composition and distribution. They identify Type 1 splash-form spherules (tektites or microtektites) that occur in the melt-ejecta (basal or lower) layer of the regional debris layer where it has a two-layered structure. These spherules are found as far from the Chicxulub site as Wyoming, but generally do not extend beyond about 4000 km away from Chicxulub. While the Type 1 particles are derived from silicic rocks, they are also mixed with sulfur-rich carbonates from the upper sediments in the Yucatan. The Type 1 spherules are poor in Ni and Ir, and the lower layer is poor in shocked quartz, consistent with their origin from the lower energy impact ejecta from the crater. Generally, the debris layer within about 4000 km of the crater is almost entirely composed of target material, rather than material from the impactor itself. Type 2 spherules, on the other hand, are found in the distal debris layer, and presumably formed primarily from the condensation of rock vapor from the impactor and target (O'Keefe and Ahrens, 1982; Johnson and Melosh, 2012b). There are subtypes of Type 2 spherules that correspond to varying composition of the original source material. Type 2 spherules occur in the upper layer in impact sites near Chicxulub, which merges into the fireball layer at distal sites. The Type 2 spherules are rich in Ni and Ir, while the fireball layer is rich in shocked quartz.

The formation of the spherical particles may depend on two different processes. Melosh and Vickery (1991) describe one formation mechanism, probably occurring in less heavily shocked portions of the target, when molten material decom-

presses until it reaches a critical line at which it starts to boil. The gas drag from the rock vapor on the molten rock spheres then tears apart the molten material, just as water droplets break apart when they fall through air. The relative velocities of water drops in air and the melt in vapor are similar, as are the surface tensions. As a result, melt droplets are similar in size to drizzle drops in light rain near 250 μm . According to Johnson and Melosh (2012b), these spherical particles are most likely to be found within 4000 km of the impact site, and to be chemically related to the target material, and not to the impactor. Such materials are reported across North America as Type 1 spherules (Bohor et al., 1987), sometimes referred to as microtektites. Since these spherules are not global, they likely were not as relevant to climate as the Type 2 spherules.

Melt droplets can also form in heavily shocked parts of the impact debris as rock vapor condenses to form melt in the fireball, which rises thousands of kilometers above the Earth's surface. These melt droplets form the Type 2 spherules. O'Keefe and Ahrens (1982) first modeled this process, and deduced that particles near a few hundred microns in size would form, as is observed. They also pointed out that the size of the spheres would be proportional to the size of the impactor. Johnson and Melosh (2012b) recently reconsidered this process for forming melt particles. They point out that the large spherules contain iridium (e.g., Smit, 1999), which is consistent with them being composed partially of the vaporized impactor. Their model of the formation and distribu-

Table 2. Gas-phase emissions (g) from the Chicxulub impact.

Sources/ gases ^d	S ($\times 10^{13}$)	C (as CO ₂ ^b) ($\times 10^{17}$)	H ₂ O ($\times 10^{15}$)	Cl ($\times 10^{12}$)	Br ($\times 10^{10}$)	I ($\times 10^7$)	N ($\times 10^{14}$)	Vertical distribution
Ambient burden (g)	1 ^a	8.4	1.3 strat	2.3 strat	3.1 strat	< 2.3 strat	2 as N ₂ O	
Impactor	4×10^3	0.3	200	7×10^2	5×10^2	7×10^4		As Type 2 spherules
Forest fires	40	6	1500	200	1000	9×10^5	10	As soot
Vaporized sea- water	60	Small	600	1×10^4	5×10^3	40	–	As Type 2 spherules
Splashed sea- water ^c	500	Small	5×10^3	1×10^5	4×10^4	3×10^2	–	Uniformly mixed above tropopause
Impact site (vaporized)	5000	0.6	90	800	400	3		As Type 2 spherules
Impact site (degassed)	500	0.1	120	2×10^3	1×10^3	7		Uniformly mixed above tropopause
Air heating							300 as NO _x created from air	Half uniformly mixed, half as Type 2 spherules

^a Based on Pinatubo eruption. ^b Mass is given in terms of C, but emission is in the form of CO₂. ^c S, Cl, Br, I likely injected as particulates. ^d The scaling factors given in parentheses apply to all values in column.

tion of these particles suggests the particles have a size that varies spatially over the plume. Averaging over the simulated plume yields a mean size of 217 μm with a standard deviation of about 47 μm for a 10 km diameter impactor hitting at 21 km s^{-1} . From the two examples given by Johnson and Melosh (2012b) it appears that the standard deviation is consistently 22 % of the mean radius for asteroids of different sizes. The initial values for the various properties of Type 2 spherules described above are summarized in Table 1 for the K-Pg impactor.

Smit (1999), who refers to the Type 2 spherules in the distal layer as microkrystites, estimated that these particles typically have a diameter near 250 μm , and a surface concentration of about 20 000 particles cm^{-2} over the Earth. Unfortunately, we are not aware of studies that measure the dispersion of the size distribution, or the spatial variation of the abundance of these particles. We assume that the particles have the density of CM2 asteroids, since Cr isotope ratios suggest that is the composition of the K-Pg impactor (Trinquier et al., 2006). Assuming this density, $\sim 2.7 \text{ g cm}^{-3}$, the mass of spherules per unit area of the Earth is about 0.4 g cm^{-2} , and the initial optical depth is about 20, as noted in Table 1. These spherules compose about half of the mass of the distal layer. We assume the particles were initially distributed uniformly around the globe, with the initial mixing ratio in the atmosphere varying only in altitude. Some theoretical studies, such as Kring and Durda (2002) and Morgan et al. (2013), suggest that these particles were not uniformly

deposited in latitude and longitude, but had focusing points such as the antipodes of the impact site. Unfortunately, we are not aware of quantitative data on the global distribution of the spherules. The study by Morgan et al. (2013) may also be more applicable to the Type 1 spherules since their numerical model does not produce vaporized material from the asteroid impact.

According to the simulations of Goldin and Melosh (2009), the infalling spherical particles reached terminal fall velocity near 70 km altitude, at which point they begin to behave like individual airborne particles. Kalashnikova et al. (2000) investigated incoming micrometeorites in the present atmosphere, which generally ablate near 85 km. Kalashnikova et al. (2000) found material entering from space stops in the atmosphere after it encountered a mass of air approximately equal to its own mass. Therefore, the altitude distribution is taken to be Gaussian, centered at 70 km and with a half width of one atmospheric scale height (about 6.6 km based on the US standard atmosphere). A scale height is chosen as the half width of the injection profile since it is a natural measure of the density of the atmosphere. Figure 1 illustrates the vertical injection profile of the spherules (green curve). As discussed below, we expect several materials with origins similar to those of the spherules to be injected in this same altitude range, but others with origins unrelated to the impact generated plume, such as soot from fires, to be injected at lower altitudes.

Table 3. The 1 km land^a injection scenario for impactor mass 1.4×10^{15} g; impactor energy $\sim 2.8 \times 10^{20}$ J = 6.8×10^4 Mt.

Property/ constituent	Type 2 spherules	Soot ^b	Nanoparticles from vapor- ized rock ^c	Clastics, < μm distributed glob- ally	S
Material amount g, column density (g cm^{-2})	1.4×10^{15} (2.6×10^{-4})	2.8×10^{13} (5.6×10^{-6})	1×10^{15} (2×10^{-4})	2.6×10^{13} (5×10^{-6})	4.4×10^{13} ($2.6 \times 10^{-5} \text{ g cm}^{-2}$ as SO ₄)
Estimated global optical depth as $1 \mu\text{m}$ particles	0.2 (as $15 \mu\text{m}$ particles)	4.7×10^{-2}	1.5	4×10^{-2}	0.22
Vertical & horizontal distributions	Table 1; over 50 % of Earth	50 % Eq. (2) + 50 % Eq. (3); over $4 \times 10^4 \text{ km}^2$	Table 1; over 50 % of Earth	Uniformly mixed above tropopause, spread over $4 \times 10^5 \text{ km}^2$	Follow nanoparticles
Optical properties	Not relevant	Table 1	Table 1	Depends on im- pact site	Table 1
Initial particle size (μm)	$15 \mu\text{m}$	Table 1	20 nm	Table 1	

^a We assume a 1 km asteroid impact would not penetrate through the 5 km average depth of the ocean. Therefore, none of the materials in this table would be injected into the atmosphere for an ocean impact. For the density of all materials, follow Table 1. ^b The material amount assumes an impact into a region where 2.25 g C cm^{-2} flammable biomass is consumed. The material amount can be scaled linearly for other choices of available biomass that burns. ^c We assume about 35 % of the impactor and an equivalent mass of target would vaporize and end up as nanoparticles. This value is an upper limit. The lower limit is zero.

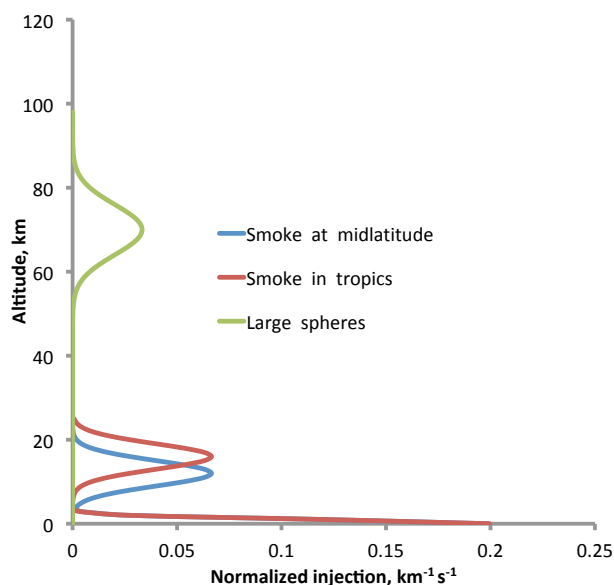


Figure 1. Injection profiles for smoke at midlatitudes and the tropics and for large spherical particles. Many other constituents follow the same vertical profiles as noted in Tables 1–4. We suggested clastics be placed above the tropopause using a constant mixing ratio.

The 70 km injection altitude refers to the level at which the large spherical particles reached terminal velocity. However, as is evident from the optical depth, many spherules

entered through the same air mass. The column mass of the distal layer is $\sim 1 \text{ g cm}^{-2}$ so the air pressure needs to about 1 hPa for the air mass above the altitude in question and the particle mass to be comparable. A pressure of 1 hPa occurs at about 48 km. Therefore, if the entire distal layer mass is placed into a model above 48 km, its mass mixing ratio will be greater than 1, and the atmosphere will be significantly out of hydrostatic balance. We are not aware of any simulations of the first few hours after the impact, but significant turbulence and mixing must have occurred as the atmosphere adjusted to the large mass imbalance. Model initialization should be checked to determine if the planned simulations start out of hydrostatic balance. If so, the injection altitude should be lowered below 70 km.

The energy release from the reentry of the large spherical particles into the atmosphere was likely responsible for setting most of the aboveground terrestrial biosphere on fire. However, due to their size, the spherules could not have remained in the atmosphere for more than a few days. Hence, they likely did not have a significant direct impact on the climate, but fell to Earth like a gentle rain.

2.1.2 Large spherules from a 1 km diameter asteroid impact

Type 1 spherules, melt droplets, will form from impacts by 1 km diameter asteroids, and produce millimeter-sized particles in the ejecta curtain layer located near the crater (John-

Table 4. Gas-phase emissions (g) from a 1 km diameter impact.

Sources/ Gases ^d	S ($\times 10^{13}$)	C ^a ($\times 10^{17}$)	H ₂ O ($\times 10^{15}$)	Cl ($\times 10^{12}$)	Br ($\times 10^{10}$)	I ($\times 10^7$)	N ($\times 10^{14}$)	Vertical distribution
Ambient burden (g)	1 ^b	8.4	1.3 strat	2.3 strat	3.1 strat	< 2.3 strat	2 as N ₂ O	
Impactor/ land only	4.4	3×10^{-2}	0.2	0.7	0.5	68	–	As Type 2 spherules
Forest fires/land only	2.7×10^{-2}	4×10^{-3}	0.9	0.12	0.62	560	6.9×10^{-3}	As soot
Vaporized seawater	0.9	small	10	200	80	0.6		Uniformly mixed
Splashed seawater ^c	3	small	30	600	200	2		
Air heating							0.6	Uniformly mixed

^a Mass is given in terms of C, but emission is in the form of CO₂. ^b Based on Pinatubo volcanic eruption. ^c S, Cl, Br, I may be released as particulates. ^d Scaling factors given in parentheses apply to all values in the column.

son and Melosh, 2014). We do not expect an impact by a 1 km diameter asteroid to create a global layer of Type 2 spherules (Toon et al., 1997). Like O’Keefe and Ahrens (1982), Johnson and Melosh (2012b) conclude that the particle size will vary in proportion to the impactor diameter and the impactor velocity. For a 1 km diameter impactor hitting the land at 20 km s^{-1} , they suggest that the mean diameter of the spherical particles will be about $15 \mu\text{m}$, with somewhat larger sizes as the impact velocity increases to 30 km s^{-1} . Table 3 provides our assumed properties of the spherules from a hypothetical 1 km diameter impactor hitting the land. It is likely that spherules would not be distributed over all of the globe for the 1 km diameter impact. Johnson and Melosh (2012a) as well as Glass and Simonson (2012) report a spherule layer associated with the Popigai impact in the late Eocene which Johnson and Bowling (2014) suggest was global in extent. This layer contains spherules similar in size or even larger than those associated with the Chicxulub impact. However, this layer is only about 10 % as thick as the distal layer from the Chicxulub impact. A 1 km impactor hitting the deep oceans may not produce a layer of spherules.

2.2 Soot

2.2.1 Soot from the Chicxulub impact

Spherical soot (also referred to as black carbon, or elemental carbon) particles were discovered in the boundary layer debris at sites including Denmark, Italy, Spain, Austria, Tunisia, Turkmenistan, the United States, and New Zealand (among others) by Wolbach et al. (1985, 1988, 1990a, b). Soot was also found in anaerobic deep-sea cores from the mid-Pacific (Wolbach et al., 2003). Soot was apparently lost by oxidation in aerobic deep-water sites in the 66 million years since emplacement. There is debate about whether these particles originated from global wildfires, or from the impact itself (Belcher et al., 2003, 2004, 2005, 2009; Belcher,

2009; Harvey et al., 2008; Robertson et al., 2013a; Pierazzo and Artemieva, 2012; Premović, 2012; Morgan et al., 2013; Kaiho et al., 2016). Robertson et al. (2013a), Pierazzo and Artemieva (2012), Premović (2012) and Morgan et al. (2013) argue that it is implausible that there was enough carbon at the impact site to produce the amount of soot observed by Wolbach et al. (1988). This debate about the origin of the particles does not greatly affect the impact these particles would have had on the climate when they were suspended in the atmosphere. The particles are small and widely distributed. They are numerous and so must have produced a very large optical depth and, being composed of carbon, they would have been excellent absorbers of sunlight. Whether the soot particles originated from global fires and were deposited in the upper troposphere, or they originated at the impact site and were deposited in the mesosphere, the climate effect of the observed soot would have been very great. Some have suggested that the soot resulted from wildfires in dead and dying trees that occurred well after the impact. However, Wolbach et al. (1988, 1990b) show that soot and iridium are tightly correlated and collocated. Indeed, Wolbach et al. (1990b) suggest the soot and iridium may have coagulated in the atmosphere. The soot and iridium in the distal layer must have been deposited within a few years of the impact, since small particles will not stay in the air much longer. Therefore, any fires must have been very close in time to the impact, and were likely contemporaneous.

Wolbach et al. (1988) estimated the global mass of elemental carbon (including aciniform soot, charcoal and any unreactive aromatic kerogen) in the debris layer as $7 \pm 4 \times 10^4 \text{ Tg}$ of C or equivalently $13 \pm 7 \text{ mg C cm}^{-2}$ based on data from five sites. Wolbach et al. (1990b) updated these mass determinations to $5.6 \pm 1.5 \times 10^4 \text{ Tg}$ or $11 \pm 3 \text{ mg C cm}^{-2}$ based on data from 11 sites. This mass of elemental carbon would require that the bulk of the above-ground biomass burned and was partially converted to el-

elemental carbon with an efficiency of about 3 %, assuming the biomass is 1.5 g C cm^{-2} of aboveground, dry organic mass per square centimeter over the land area of Earth. This biomass density is typical of current tropical forests. This inferred 3 % emission factor is about 60 times greater than that suggested by Andreae and Merlet (2001) for current wildfires, but agrees with laboratory and other observations from burning wood under conditions consistent with mass fires (Crutzen et al., 1984; Turco et al., 1990). Mass fires are more intense than forest fires, and consume all the fuel available, possibly including that in the near-surface soil. Ivany and Salawitch (1993) argued independently from oceanic carbon isotope ratios that at least 25 % of the aboveground biomass must have burned at the K-Pg boundary.

Wolbach et al. (1990b) distinguish several forms of elemental carbon. Aciniform carbon is composed of grape-like clusters of 0.01 to 0.1 μm spherules. On average, this type of soot is 26.6 % of the elemental carbon, yielding a global mass abundance of $1.5 \times 10^4 \text{ Tg}$ of aciniform carbon. Charcoal is estimated at 3.3 to $4.1 \times 10^4 \text{ Tg}$, and unreactive kerogen at 0 to $0.8 \times 10^4 \text{ Tg}$. Wolbach et al. (2003) discuss a data set from the mid-Pacific that suggests aciniform soot is $9 \times 10^3 \text{ Tg}$, and charcoal is also $9 \times 10^3 \text{ Tg}$. Wolbach et al. directly measure the carbon content of their samples. The aciniform soot to charcoal ratio is determined by using an electron microscope to distinguish small and large particles.

There are several uncertainties in determining the amount of soot to use in a model. An upper limit of the amount injected into the stratosphere is $7.1 \times 10^4 \text{ Tg}$ based on the upper error bar of the Wolbach et al. (1990b) elemental carbon values. An important assumption in this upper limit is that the larger particles found by Wolbach et al. (1990b), are either aggregates of smaller ones, or of the same general size as the aggregates of the smaller ones that occur after coagulation. A lower limit of $1.1 \times 10^4 \text{ Tg}$ is obtained using the lower error bar of the elemental carbon from Wolbach et al. (1990b), and assuming 26.6 % is aciniform soot. Alternatively, one could argue that this lower limit of aciniform soot should be injected into the stratosphere, along with $3.3 \times 10^4 \text{ Tg}$ of charcoal using different size distributions. The most likely value of the aciniform soot in the stratosphere is $1.5 \times 10^4 \text{ Tg}$, and of elemental carbon $5.6 \times 10^4 \text{ Tg}$. We use these most likely values in Table 1.

Kaiho et al. (2016) argue that the soot came from burning hydrocarbons in the crater and that the total mass emitted was either 5×10^2 , 15×10^2 , or $26 \times 10^2 \text{ Tg}$. If we reduce these values by the authors' factor of 2.6 to represent the stratospheric emissions, they are 0.4, 1.0, and 1.7 % of the globally distributed elemental carbon reported by Wolbach et al. (1990b).

Kaiho et al. (2016) measured several polycyclic aromatic hydrocarbons (PAHs) that are minor components of soot from one distal site in Caravaca de la Cruz, Spain, and another site at Beloc, Haiti, that is about 700 km from the crater. Since the PAHs measured are minor constituents of soot,

Kaiho et al. (2016) need to use a large correction factor to determine the amount of soot. They first multiply by factors of 2, 5.9, or 10 to account for possible loss of PAH concentrations over time. They present no data to justify these factors. They then multiply by 3.3×10^3 citing this as the ratio of their measured PAHs to soot in diesel soot. No error bars were presented for this factor, and no values were given for the ratio in biomass soot. The origin of this correction factor is not evident in the cited reference. They then multiplied by another factor of 2.6 to represent the fraction of their soot estimate that they suspect reached the stratosphere. Their overall correction factors were therefore 17×10^3 , 50×10^3 , and 86×10^3 . Given these large correction factors, and the lack of information about their uncertainty, it is difficult to compare them with the direct determinations done by Wolbach et al. (1990b), which do not require any correction factors.

As noted in Table 1, the mass of soot found by Wolbach et al. (1988) would produce an optical depth near 100 if the particles coagulated to spheres with a radius of 1 μm while they were in the atmosphere. Toon et al. (1997) pointed out that soot clouds with such a large optical depth would reduce light levels at the Earth's surface effectively to zero. The optical and chemical evolution of the particles once in the atmosphere may be influenced by the presence of liquid organics on the soot particles. Bare soot particles coagulate into chains and sheets, while particles that are coated by liquids may form balls. Chains, sheets, and coated balls have very different optical properties than spheres (Wolf and Toon, 2010; Ackerman and Toon, 1981; Bond and Bergstrom, 2006; Mikhailov et al., 2006). Particulate organic matter can be absorbing, and soot coated with organics can have enhanced absorption relative to soot that is uncoated (Lack et al., 2012; Mikhailov et al., 2006). These fractal shapes and organic coatings might not be preserved in samples in the distal layer since all the particles have been consolidated in a layer, and even in the current atmosphere the organics have short lifetimes due to rapid oxidation.

Wolbach et al. (1985) fit the size of the particles they observed, after exposing them to ultrasound to break up agglomerates, to a lognormal size distribution, described by

$$\frac{dN}{d \ln r} = \frac{N_t}{\ln \sigma \sqrt{2\pi}} \exp \left[- \left(\ln^2 \left(\frac{r}{r_m} \right) / 2 \ln^2 \sigma \right) \right]. \quad (1)$$

Here, r is the particle radius, N_t is the total number of particles per unit volume of air, r_m is the mode radius, and σ is the width of the distribution. Wolbach et al. (1985) found $r_m = 0.11 \mu\text{m}$ and $\sigma = 1.6$ for the soot in the K-Pg boundary layer. We assume this distribution represents the initial sizes of the soot particles. The final size, which would be determined by coagulation while in the atmosphere, might not be preserved in the sediments, and loosely bound clumps of particles would have been destroyed by the ultrasound treatment of the samples.

The size distribution of soot from the K-Pg boundary is similar to that of smoke near present-day biomass fires as

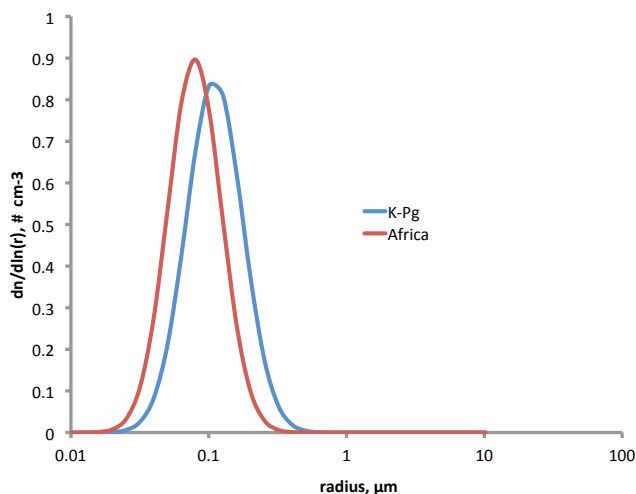


Figure 2. The size distributions for smoke from modern fires in Africa and from the K-Pg boundary layer (Wolbach et al., 1985; Matichuk et al., 2008).

indicated in Fig. 2 (e.g., Matichuk et al., 2008). This similarity in sizes is somewhat surprising because the present-day smoke size distribution includes organic carbon, which is present in addition to the elemental carbon (soot). Generally, in wildfire smoke, organic carbon has 5–10 times the mass of soot, so one might anticipate that the K-Pg soot would be about half the size of the present-day smoke rather than of similar size since the organic coatings are no longer present, or were never present, on the K-Pg soot. The organics might never have been present because mass fires are very intense and tend to consume all the available fuel, which might include the organic coatings. Aggregation in the hot fires may have caused this slightly larger than expected size in the K-Pg sediments. Wolbach et al. (1985) suspended their samples in water and subjected them to ultrasound for 15 min in a failed attempt to completely break up agglomerates. This failure indicated that the remaining agglomerates might have been flame welded. Therefore, the K-Pg size distribution from Wolbach et al. (1985) does not represent the monomers in the aggregate soot fractal structures. Rather the K-Pg size distributions represent a combination of monomers and aggregates that may have formed at high temperatures. Possibly the smallest-sized particles measured by Wolbach et al. (1985), which have radii of 30–60 nm, represent the soot monomers. These are in the same general range as monomer sizes observed in soot from conventional fires (Bond and Bergstrom, 2006).

The injection altitude of the soot depends on its source. In a series of papers, Belcher et al. (2003, 2004, 2005, 2009) and Belcher (2009) argue from multiple points of view that there were no global forest fires. Harvey et al. (2008) and Kaiho et al. (2016) argue that the soot originated from oil, coal, and other organic deposits at the location of the impact. If correct, the soot might have been injected at high altitude along

with the large spherules. Recently, Robertson et al. (2013a) reconsidered each of the arguments presented by Belcher et al. (2003, 2004, 2005, 2009) and Belcher (2009) and came to the conclusion that global wildfires did indeed occur. Pierazzo and Artemieva (2012), Premović (2012), Morgan et al. (2013), as well as Robertson et al. (2013a) have independently argued that oil and other biomass in the crater is quantitatively insufficient to be the source of the soot. Therefore, we assume that the soot indeed originated from burning biomass distributed over the globe. The soot is clearly present in the distal layer material, and therefore was once in the atmosphere where it could cause significant changes to the climate.

Toon et al. (2007) have outlined the altitudes where one expects large mass fires to inject their smoke. Numerical simulations have shown that mass fires larger than about 5 km in diameter have smoke cloud tops well into the stratosphere. The smoke itself is distributed over a range of heights, however. The details of the injection profiles depend on the rate of fuel burning, the size of the fires, and the meteorological conditions among other factors. In addition, some smoke is quickly removed from the atmosphere by precipitation in pyrocumulus. However, it is thought that overseeding of the clouds by smoke prevents precipitation, and that only 20 % or so of the smoke injected into the upper troposphere is promptly rained out (Toon et al., 2007). Smoke that is injected near the ground, on the other hand, will be removed by rainfall within days or weeks.

The K-Pg impact occurred at a time when average biomass density likely was higher than now. Following Small and Heikes (1988; their Fig. 3f) and Pittcock et al. (1989) one would expect smoke from large-area fires burning in high biomass density areas to show a bi-modal smoke injection profile. The smoke at higher levels is injected in the pyrocumulus and other regions with strong vertical motions. However, once the fires die-down smoke will be emitted in the boundary layer. There are also downdrafts, as well as entrainment and mixing with the environment, that occur in all cumulus and these will carry some smoke into the boundary layer. We simulate this with injections whose vertical distributions are Gaussian functions centered at the tropopause and at the surface, as illustrated in Fig. 1. The injection at the tropopause (Eq. 2) has a half width of 3 km, but nothing is injected above about 25 km. We set this upper altitude limit based on the heights of the stratospheric sulfate clouds from explosive volcanic eruptions, which rise buoyantly as do smoke plumes. The Gaussian distribution at the ground (Eq. 3) has a half width of 1 km, assuming that the local boundary layer is relatively shallow. We assume 50 % of the soot is contained in each of these distributions (Eqs. 2 and 3) for the general case, and for the 1 km impact. For the K-Pg, we assume the soot observed in the distal layer by Wolbach et al. (1988, 1990b) was all in the portion of the Gaussian distribution at the tropopause (Eq. 2).

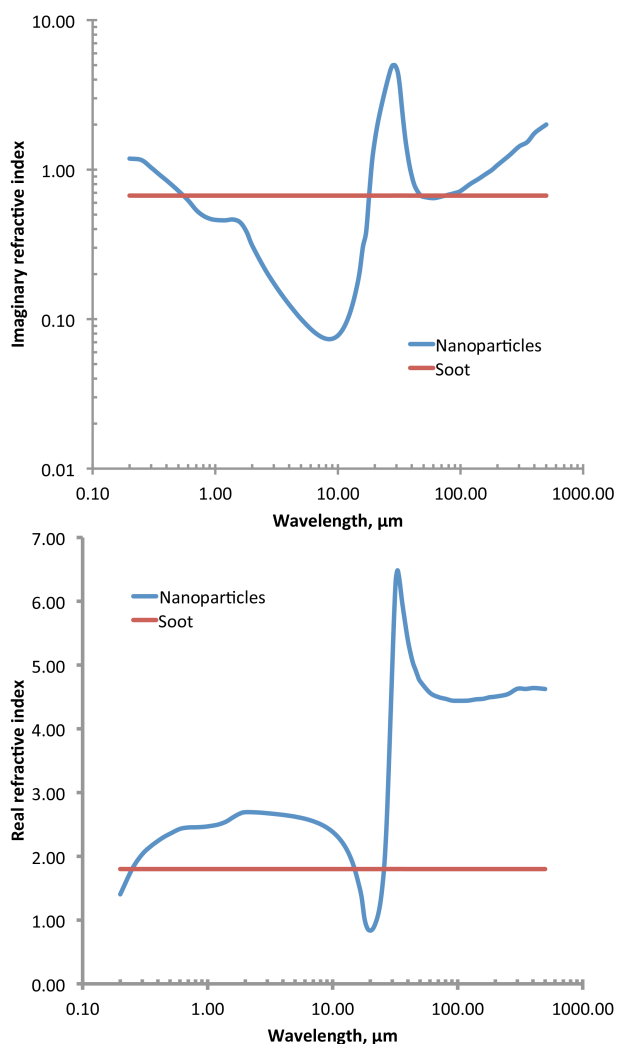


Figure 3. The real and imaginary parts of the refractive index suggested for nanoparticles, and for soot.

Therefore, the injection profiles are given by

$$I(\text{g s}^{-1} \text{ km}^{-1}) = \frac{I_{T1}}{\eta\sqrt{2\pi}} \left[e^{\left(-0.5\left(\frac{z-z_{\text{trop}}}{\eta}\right)^2\right)} \right] \quad (2)$$

$$I(\text{g s}^{-1} \text{ km}^{-1}) = \frac{I_{T2}}{\mu\sqrt{2\pi}} \left[e^{\left(-0.5\left(\frac{z}{\mu}\right)^2\right)} \right] \quad (3)$$

Here, I is the mass emission rate per kilometer of altitude, I_{T1} and I_{T2} are the total mass emitted per second into the upper (Eq. 2) or lower (Eq. 3) altitude range after correcting for the emission altitude range (0–25 km) and grid spacing, μ is 1 km, η is 3 km, and z_{trop} is the altitude of the tropopause.

Geographically, we assume for the K-Pg event that all the surface biomass is set on fire. For the 1 km diameter impact, however, only the region near the impact site would burn as discussed further below.

There is also an issue of how long it takes to inject the smoke. Forest fires often burn for days, advancing along a fire front as winds blow embers far beyond the flames and onto unburned terrain. Mass fires may not spread because powerful converging winds restrict the spread. However, little is known observationally about mass fires, and fires can spread by intense infrared radiation lighting adjacent material. If mass fires are restricted then they will burn only as long as they have fuel. The present aboveground global biomass in tropical forests is in the range of 0.6–1.2 g C cm⁻² (Houghton, 2005). The energy content of the biomass is on the order of 3 × 10⁴ J (gC)⁻¹ or, given the biomass concentration just mentioned, about 3 × 10⁸ J m⁻². Penner et al. (1986) and Small and Heikes (1988) found that large area mass fires with energy release rates of 0.1 MW m⁻² would have plumes reaching the lower stratosphere. Hence, it would be necessary to assume that the fuel burned in an hour or so to achieve these energy releases. Of course, it might take some time for fires in different places to start fully burning, so considering the entire region of the mass fire, as opposed to a small individual part of the fires, might prolong the energy release considerably. For example, it took several hours for the mass fire in Hiroshima to develop after the explosion of the atom bomb (Toon et al., 2007).

It should be noted that in simulations of stratospheric injections of soot from nuclear conflicts, soot is self-lofted by sunlight heating the smoke (Robock et al., 2007b). However, in the case of the K-Pg impact, if there are other types of particles injected above the soot, which then block sunlight, the soot may not be self-lofted, which will limit its lifetime. The initial soot distribution that is estimated here does not include the effects of self-lofting, which would continue after the initial injection and should be part of the climate simulation.

The final property to specify for soot is the optical constants. This issue is complicated by the possible presence of organic material on the soot (Lack et al., 2012). However, it is known that many of these organics are quickly oxidized by ozone, which is plentiful in the ambient stratosphere. The stratosphere after the impact however, may have become depleted in ozone very quickly, so that the organic coatings might have survived. It is also possible that intense fires, such as mass fires, will consume the organic coatings, which may explain why the production of soot in the fires seems to have been so much more efficient than for normal fires. It may therefore be sufficient to treat the soot as fractal agglomerates of elemental carbon (Bond and Bergstrom, 2006). It is known that the optical properties of the agglomerates will not obey Mie theory. However, one may treat their optical properties as well as their microphysical properties using the fractal optics approach described by Wolf and Toon (2010). The optical constants for elemental carbon may then be used for the monomers. Alternatively, one may add the organic mass to the particles, and treat them using core-shell theory (Toon and Ackerman, 1981; Mikhailov et al., 2006).

Bond and Bergstrom (2006) have thoroughly reviewed the literature on the optical properties of elemental carbon. They conclude that the optical constants are most likely independent of wavelength across the visible, with a value that depends on the bulk density of the particles. Following their range of values for refractive index vs. particle density, we suggest using a wavelength-independent real index of refraction $n = 1.80$ and an imaginary index $k = 0.67$. We also use these values in the infrared as shown in Fig. 3. For the monomers in Tables 1 and 3, we adopt the density suggested by Bond and Bergstrom (2006) for light-absorbing material, 1.8 g cm^{-3} .

2.2.2 Soot from a 1 km impact

Extrapolations of the soot injection parameters to smaller impactors than the one defining the K-Pg boundary should only involve changes to the mass of soot injected, since the basic properties of the soot at the K-Pg boundary are similar to those of forest fire soot. Therefore, the particle sizes, injection heights, and optical constants recommended in Table 3 for the smaller impact are the same as listed in Table 1 for the Chicxulub impact. The mass of soot injected is estimated from the extrapolations in Toon et al. (1997). For an impactor as small as 1 km diameter, debris from the impact site would not provide sufficient energy to ignite the global biota since the energy of the 1 km impactor is about 1000 times less than that of the Chicxulub impactor. Instead, radiation from the ablation of the incoming object and from the rising fireball at the impact site would ignite material that is within visible range of the entering object and the fireball. This ignition mechanism is well understood from nuclear weapons tests (Turco et al., 1990). Hence, for a 1 km diameter impactor the fuel load at the site of the impact becomes critical to evaluate the soot release. No soot would be produced from an impact in the ocean, an ice sheet, or a desert. In Table 3, to compute the smoke emitted (28 Tg), we use Eq. (12) from Toon et al. (1997) to obtain an area of $4.1 \times 10^4 \text{ km}^2$ for the expected area exposed to high thermal radiation density from the fireball for a 1 km diameter impactor with an assumed energy of $6.8 \times 10^4 \text{ Mt}$. We then multiply that area by 3 % (the fraction of C in the burned fuel that is converted to smoke) and by 2.25 g C cm^{-2} (the assumed carbon content per unit area of the dry biomass that burns). The user of Table 3 can choose alternate values of the injected soot by scaling linearly to the biomass concentration they chose.

Ivany and Salawitch (1993) suggest that the land average aboveground biomass was about $1 \times 10^{18} \text{ g}$ (about 0.7 g C cm^{-2}) at the end of the Cretaceous. The current land average, aboveground biomass is about 0.3 to 0.44 g C cm^{-2} (Ciais et al., 2013). An additional 1 to 1.6 g C cm^{-2} is currently present in the soil, while Ivany and Salawitch suggest 1 g C cm^{-2} in the soil in the Cretaceous. Some of the soil biomass may burn in a mass fire. Tropical and boreal forests currently have average biomass concentrations (aboveground

and in soil) of about 2.4 g C cm^{-2} , while temperate forests have about 1.6 g C cm^{-2} , including soil carbon (Pan et al., 2011). Soil carbon is 30 % of carbon in tropical forests and 60 % in boreal forests. Together, tropical and boreal forests cover 6 % of the Earth's surface, and temperate forests 1.5 %. These forests cover 26 % of Earth's land area. In Table 3, we assume that the biomass that burns is typical of a tropical or boreal forest, assuming the soil carbon burns. The reader can make other choices for the biomass by scaling from the fuel load that the reader prefers.

Another modeling issue of concern is the ability of models to follow the initial evolution of the plume. If we assume that half of the 28 Mt of smoke from the 1 km impact is injected over an area of $4 \times 10^4 \text{ km}^2$ and over a depth of 6 km near the tropopause (Eq. 2) as $0.1 \mu\text{m}$ radius smoke particles, the smoke will have an initial optical depth near 4000, and the number density of particles will be about 10^7 cm^{-3} . (The other half of the smoke mass injected near the ground (Eq. 3) will likely be removed quickly and have little impact on climate.) Intense solar heating at the top of the smoke cloud near the tropopause will loft it, while coagulation will reduce the number of particles by a factor of 2 and increase their size proportionately in only 1 min. Hence, one needs to model this evolution on sub-minute time scales to accurately follow the initial evolution. Alternatively, but less accurately, one might spread out the injection in time and space, so that the climate model can track the evolving smoke cloud using typical model time steps.

2.3 Nanoparticles from vaporized impactors

2.3.1 Nanoparticles from the vaporized material following the Chicxulub impact

Johnson and Melosh (2012b) found that, at the end of their simulations of the rising fireball, about 44 % of the rock vapor that was created from the K-Pg asteroid impact remained as vapor rather than condensing to form large spherules. This vapor is about an equal mixture of impactor and asteroid, so the 44 % mass fraction is approximately equal to the mass of the impactor. This 44 % vapor fraction depends on the pressures reached in the impact, the equation of state of the materials, as well as the detailed evolution of the debris in the fireball. The fate of this vapor-phase material is not well understood and has been little studied. It may simply have condensed on the spherules, or it may have remained as vapor.

Presently, $100 \mu\text{m}$ and larger-sized micrometeoroids ablate to vapor in the upper atmosphere. Hunten et al. (1980), following earlier suggestions, modeled the condensation of these rock vapors as they form nanometer-sized particles in the mesosphere and stratosphere. Bardeen et al. (2008) produced modern models of their distribution based on injection calculations from Kalashnikova et al. (2000). Hervig et al. (2009) and Neely et al. (2011) showed that these tiny particles are observed as they deposit about 40 t of very fine-

grained material on Earth's surface per day. It is possible that a similar process occurred after the Chicxulub impact. However, in the Chicxulub case the vaporization occurred during the initial asteroid impact at Chicxulub rather than on reentry of the material after the fireball rose thousands of kilometers into space and dispersed over the globe.

The presence of 15–25 nm diameter iron-rich material has been recognized in the fireball layer at a variety of sites by Wdowiak et al. (2001), Verma et al. (2002), Bhandari et al. (2002), Ferrow et al. (2011), and Vajda et al. (2015), among others. The nanophase iron correlates with iridium and is found worldwide, and therefore is likely a product of the impact process. Unfortunately, these authors have not quantified the amount of this material that is present. Berndt et al. (2011) were able to perform very high-resolution chemical analyses, and also report a component of the platinum group elements that arrived later than the bulk of the ejecta, and was probably the result of submicron-sized particles. However, they were not able to size the particles, nor quantify their abundance.

In Table 1, we take the upper limit of the injected mass of nanoparticles to be 2×10^{18} g. The lower limit is zero. This choice for the upper limit is consistent with the vapor mass left at the end of the simulations by Johnson and Melosh (2012b). We assume an initial diameter of 20 nm, following Wdowiak et al. (2001). We assume the particles are initially injected over the same altitude range as the Type 2 spherules, because we speculate that the small particles would not separate from the bulk of the ejecta in the fireball until the ejecta entered the atmosphere and reached terminal velocity. The mass injected would lead to an optical depth of particles larger than 1000 even if they coagulated into the 1 μm size range. Goldin and Melosh (2009) point out that such an optically thick layer of small particles left behind by the falling large spheres might also be important for determining whether the infrared radiation from the atmosphere heated by the Type 2 spherules is sufficient to start large-scale fires.

The optical properties of the nanoparticles are not known. We suggest using the optical properties of the small vaporized particles currently entering the atmosphere from Hervig et al. (2009). These optical constants are plotted in Fig. 3. We also assume that the particles have the density of CM2 asteroids, since Cr isotope ratios suggest that is the composition of the K-Pg impactor (Trinquier et al., 2006). This density is 2.7 g cm^{-3} . A significant fraction of the vaporized material may be from the impact site, so using an asteroidal composition to determine the density is an approximation.

2.3.2 Nanoparticles from the vaporized material from a 1 km impact

Johnson and Melosh (2012b) did not comment on the amount of vapor that would be expected to not condense as spherules from a 1 km diameter impact. From the theory of impacts,

it is expected that an amount of impactor plus target that is about twice the mass of the impactor would be converted into vapor from a 1 km diameter impact, just as it is for a 10 km diameter impact. In Table 3, we assume that as an upper limit 35 % of the impactor mass plus an equivalent amount of target material would be left as vapor after spherules form. We chose this mass fraction, which is lower than that for the K-Pg object, because the 1 km impact will have a smaller fireball, and be more confined by the atmosphere. We also assume the injected particles will have a diameter of 20 nm. From simple energy balance along a ballistic trajectory, we would expect that the vaporized ejecta in the fireball from a 1 km impact would rise about a thousand kilometers above the Earth's surface. This altitude is consistent with limited numerical calculations for large energy releases, which indicate that the vertical velocity of the fireball is not significantly reduced in passing through the atmosphere (Jones and Kodis, 1982). As the material reenters the atmosphere, the particles will come to rest when they encounter an atmospheric mass comparable to their own mass. Hence, it is likely that the altitude distribution of the nanoparticles from the 1 km impact will be the same as we have assumed for the K-Pg impactor in Table 1, which is also similar to but slightly lower in altitude than the vertical distribution of micrometeorites on present-day Earth, as discussed by Bardeen et al. (2008). It is difficult to determine precisely the area that will be covered by this material as it reenters the atmosphere. If we assume that it takes about 30 min for the debris to reach peak altitude and return to the Earth and that the plume is spreading horizontally at about 4 km s^{-1} then the debris would enter the atmosphere over an area of about half that of the Earth. These estimates of area covered are consistent with the observations of the SL-9 impact collisions with Jupiter, and the plume from the much less energetic impact at Tunguska, though these are not perfect analogs (Boslough and Crawford, 1997). The optical depth of the nanoparticles from the 1 km diameter impact averaged over the Earth is estimated for comparison with the estimates of other types of particles to be relatively large (1.5) as noted in Table 3.

2.4 Submicron clastics

2.4.1 Submicron clastics from the Chicxulub impact

Another clear component of the K-Pg debris layer is pulverized target material. This clastic material was first recognized from shocked quartz grains (Bohor, 1990), but there are also shocked carbonate particles from the Yucatan Peninsula in the K-Pg boundary layer material (Yancy and Guillemette, 2008; Schulte et al., 2008). Because of chemical alteration of much of this material in the past 65 million years, it is difficult to determine the mass and size distribution directly except for the shocked quartz, which is readily identified. The shocked quartz grains generally are large and would not have remained long in the atmosphere. However, the shocked

quartz is probably not directly related to the bulk of the clastics. For instance, within 4000 km of Chicxulub, the shocked quartz is primarily in the few millimeter-thick fireball layer, which is distinct from the several centimeter or thicker ejecta layer that is dominated by clastics. The shocked quartz likely came from basement rock, reached higher shock pressures than the bulk of the pulverized ejecta, and therefore was distributed globally in the impact fireball along with the melted and vaporized material from the target and impactor. The other pulverized material, in contrast, came mainly from the upper portions of the target along with basement rocks toward the exterior of the crater, and the fragments were distributed locally (within about 4000 km of Chicxulub) in the impact ejecta debris.

The submicron fraction of the clastics is of interest because particles of such size might remain in the atmosphere for months or years and perturb the climate, unlike larger particles that would be removed quickly by sedimentation. For instance, Pueschel et al. (1994) found 3–8 months after the 1991 eruption of Mt. Pinatubo in the Philippines that volcanic dust particles with a mean diameter near 1.5 μm were optically important in the lower stratosphere in the Arctic.

The optical constants for the injected clastics are suggested from their composition. For the Chicxulub impact, the clastic material is largely carbonate evaporates. We suggest using the optical constants of limestone from Orofino et al. (1998). Unfortunately, the values need to be generated from a table of oscillator strengths. They also need to be interpolated into the visible wavelength range. We suggest extending the oscillator predictions into the visible range as done by Querry et al. (1978). The density of limestone is in the range of 2.1–2.6 g cm^{-3} , while dolomite and anhydrite have densities near 2.9 g cm^{-3} . Granite has a density near 2.6–2.8. While each of these materials contribute to the clastic debris, for convenience, we assume the pulverized ejecta have a density of 2.7 g cm^{-3} .

Pope (2002) and Toon et al. (1997) used two different methods to determine the amount of the submicron-clastic material from the Chicxulub impact. Unfortunately, these estimates disagree by about 4 orders of magnitude, as indicated in Table 5, third row, columns 1 and 2. Toon et al. (1997) used arguments based mainly on impact models to estimate that more than 10 % of the mass of the distal layer ($> 7 \times 10^{17}$ g) is submicron diameter clastics, which would be significant to climate. Pope (2002) estimated that the clastics in the distal layer have a mass that is $< 10^{14}$ g. Pope (2002) used data on shocked quartz to constrain the amount of clastics, which in principle is a better approach than using estimates based on a model as in Toon et al. (1997). The amount of clastics of all sizes in the Pope (2002) model (10^{16} g) is only 12–30 times larger than the clastics of all sizes emitted in the relatively small 1980 Mt. St. Helens eruption. Therefore, based on the Pope (2002) analysis, the submicron fraction would not be of significance to climate. Below, we attempt to reconcile these

two approaches to better determine the amount of submicron clastics.

Potential errors in the Toon et al. (1997) estimate of submicron clastics

Toon et al. (1997) estimated the amount of submicron clastics starting from analytical models of the mass of material injected into the atmosphere by a 45° impact. They estimated the mass of melt plus vapor per megaton of impact energy ($\sim 0.2 \text{ Tg Mt}^{-1}$) and the mass of pulverized material per megaton of impact energy (about 4.5 Tg Mt^{-1}). Assuming a 1.5 $\times 10^8$ Mt impact, these formulae suggest a melt plus vapor amount of 3×10^{19} g ($\sim 1 \times 10^4 \text{ km}^3$, assuming a density of 2.7 g cm^{-3}) and a pulverized amount of 7×10^{20} g ($\sim 2.5 \times 10^5 \text{ km}^3$). While sophisticated impact calculations generally agree with the amount of melt plus vapor, not all of it is found to reach high enough velocity to be ejected from the crater. For example, Artemieva and Morgan (2009) investigated a number of impact scenarios that created transient craters with diameters of 90–100 km, which they thought to be consistent with the transient diameter of the Chicxulub crater. Considering those cases with oblique impacts from 30–45° with energies of 1.5–2 $\times 10^8$ Mt, they found that the melt was in the range of 2.6×10^4 to $3.8 \times 10^4 \text{ km}^3$. However, the amount that reached high enough speed to be ejected from the crater was in the range of 5×10^3 to $6 \times 10^3 \text{ km}^3$ (average 5.6 $\times 10^3 \text{ km}^3$, 1.4 $\times 10^{19}$ g, about 2–10 impactor masses). On average, only about 20 % of the melt and vapor amount escapes from the crater. Therefore, Toon et al. (1997) may have overestimated the amount of melt escaping from the crater by about a factor of 2. It should be noted that in Artemieva and Morgan (2009) the melt exceeds the mass of the distal layer, which is about 4×10^{18} g, by about a factor of 5 because much of the melt is deposited as part of the ejecta curtain and never reaches the distal region.

Artemieva and Morgan (2009) found that the total mass ejected from the crater is 1.3 $\times 10^4 \text{ km}^3$ (2.9×10^{19} g). Assuming that 90 % of this material is pulverized rock, their results imply that Toon et al. (2007) overestimated the amount of clastic debris ejected from the crater by a factor of about 25. In column 3 of Table 5, we correct the amount of pulverized material to agree with the Artemieva and Morgan (2009) value of 2.9×10^{19} g of clastics escaping the crater. It is interesting to note that the clastic mass from Chicxulub is only a factor of about 10 larger than the minimal estimated mass of clastics ejected in the Toba volcanic eruption about 70 000 years ago (Matthews et al., 2012).

Another issue is the fraction of the pulverized debris that is submicron. Toon et al. (1997) computed the amount of pulverized debris whose diameter is smaller than 1 μm from size distributions measured in nuclear debris clouds originating from nuclear tests that were many orders of magnitude lower in energy than the K-Pg impact, and from impact crater studies cited by O’Keefe and Ahrens (1982) based on grain

Table 5. Comparison of Toon et al. (1997) and Pope (2002) estimates of submicron clastics.

Method	Quartz based estimate – Pope (2002)	Injected mass – Toon et al. (1997) ^a	Injected mass – revised	Quartz based estimate – revised	1 km impactor ^b
Initial clastic debris, g	$< 10^{16}$	7×10^{20}	2.9×10^{19}	5×10^{18}	1.3×10^{16}
% clastic $< 1 \mu\text{m}$	< 1	0.1	2	1	2
Submicron clastics, g	$< 10^{14}$	7×10^{17}	5.8×10^{17}	5×10^{16}	2.6×10^{14}
Stratospheric sub-micron surviving initial removal, g	10^{14}	7×10^{17}	$< 5.8 \times 10^{16}$	5×10^{16}	$< 2.6 \times 10^{13}$

^a Assuming an impact energy of 1.5×10^8 Mt and a velocity of 20 km s^{-1} . ^b Scaled from injected mass revised using energy scaling assuming an impact energy of 6.8×10^4 Mt.

size measurements from craters. Toon et al. (1997) assume that 0.1 % of the total clastic material would be submicron. Pope (2002) cited studies of volcanic clouds to conclude that 1 % by mass of the pulverized material would be submicron.

Rose and Durant (2009) examined the total grain size distribution (TGSD) from a number of volcanic eruptions and concluded that the amount of fine ash is related to increasing explosivity of the event. The TGSD is supposed to represent the size distribution as the clastics left the crater. Mt. St. Helens is the most likely of the volcanic eruptions they considered to be relevant to the extreme energy release in a large impact. About 2 % of the total ejecta from Mt. St. Helens had a diameter smaller than $1 \mu\text{m}$. Since the erupted mass was about $3\text{--}8 \times 10^{14}$ g, the submicron mass emitted by Mt. St. Helens was about $6\text{--}16 \times 10^{12}$ g. Matthews et al. (2012) considered the Toba eruption, whose clastics are within an order of magnitude of those from Chicxulub. Their data show that 1–2 % of the mass of the clastics is in particles smaller than $1 \mu\text{m}$ and 2–6 % in clastics smaller than $2.5 \mu\text{m}$.

In Table 5, we use 2 % of the pulverized material as a revised estimate for the fraction of the clastic material that is released as submicron ejecta. This fraction is a factor of 20 larger than the one used in Toon et al. (1997). Hence, our revised submicron mass estimate for the Chicxulub impact (column 3 row 3) is very similar to the one Toon et al. (2007) estimated (column 2 row 3) because, although we lowered the estimate of the clastic mass exiting the crater to agree with Artemieva and Morgan (2009), we increased the estimate of the fraction that is submicron.

A confounding issue is the amount of submicron and other clastics that escapes from the near-crater region and is distributed globally. A large fraction of the pulverized debris in the ejecta curtain was removed within 4000 km of the impact crater (Bohor and Glass, 1995), and volcanic ejecta is likewise largely removed near the volcanic caldera. For example, there is 4–8 cm of ash 3000 km from the Toba crater, which is

not too different from the thickness of the Chicxulub deposits at a similar distance from the crater. If the removal occurred only by individual particle sedimentation, one could simply take the mass in the smaller ranges of the size distribution and assume it spread to the rest of the globe. However, it is clear from volcanic eruption data that a significant fraction of the submicron debris is removed near the volcano by processes other than direct sedimentation (Durant et al., 2009; Rose and Durant, 2009). These processes include rainout of material from water that condenses in the volcanic plume, and also agglomeration possibly enhanced by electrical charges on the particles. It is likewise clear that such localized removal occurred after the K-Pg impact. Yancy and Guillemette (2008) describe accretionary particles that make up a large fraction of the debris layer as far as 2500 km from the Chicxulub crater. These agglomerated particles, which range in size from tens to hundreds of micrometers, are composed mainly of particles with a radius of 1–4 μm . While largely composed of carbonate, the particles are enriched in sulfur.

One can use the size distributions from volcanic data, along with the total clastic mass ejected from Chicxulub to compute the particle agglomeration, and thereby follow the particles as they spread across the Earth. Such work is now being done for volcanic events, for example, by Folch et al. (2010). They found that they can successfully reproduce mass deposited on the surface from the Mt. St. Helens eruption by including agglomeration. However, such calculations for Chicxulub are difficult for several reasons: the large clastic masses involved exceed the mass of the atmosphere for a considerable distance from the crater, so the debris flows cannot be reproduced in standard climate models; the complexity of the distribution of material in the plume with some material reaching escape velocity and other parts being hurled over a substantial fraction of the planet make it difficult to determine the spatial distribution of the material, and some material is likely lofted well above the tops of most climate

models; and the presence of clastics, melt, and rock vapor together with sulfur and water produces a chemically complex plume.

Eventually, it will be necessary to use detailed nonhydrostatic, multiphase plume models including agglomeration to better understand the distribution of Chicxulub ejecta. In the meantime, for climate modeling, we suggest placing the clastic mass in Table 5 (2.9×10^{19} g) in a circular area with radius of 4000 km, which is 22.4 % of the area of Earth. This will result in a column density of 25 g cm^{-2} , or a layer thickness of about 10 cm. The mass density of the atmosphere is about 1000 g cm^{-2} , so this is about a 2.5 % perturbation to the mass of the atmosphere. In reality the mass is concentrated near the crater as shown by Hildebrand (1993). However, the observed mass density is relatively constant between 1000 and 4000 km. The initial vertical distribution of this material may be very complex due to density flows within several hundred kilometers of the crater. We suggest initializing models assuming an injection with an altitude independent mass mixing ratio of about 2.5 %. Given our suggested vertical distribution, 90 % of the material will initially lie in the troposphere. Tropospheric material is unlikely to become globally distributed even if it escapes agglomeration, because it will quickly be removed by rainfall.

As an alternative to the complexity of modeling the loss of this material in the troposphere and considering the entire size distribution, we suggest simply placing an appropriate mass into the stratosphere. The values for a stratospheric injection are given in the bottom row of Table 5 and the first row of Table 1. For illustration, we have estimated the final optical depth assuming that 10 % of the submicron material (the amount placed into the stratosphere) will escape removal. For a size distribution, we suggest using the smaller size mode measured in the stratosphere after the Mt. St. Helens eruption, as summarized by Turco et al. (1983). This size distribution is log-normal (Eq. 1), with a mode radius of $0.5 \mu\text{m}$ and a standard deviation of 1.65. The estimated optical depth of 88 is very large, even though the submicron clastic material in this estimate is only about 1 % of the mass of the distal layer.

Potential errors in the Pope (2002) estimate of submicron clastics

Pope (2002) determined the amount of clastics by modeling the amount of quartz in the distal layer. He found that he needed an initial injection of about 5×10^{15} g of quartz to match the distribution of quartz mass with distance from the impact site. It is not clear how good this estimate is because the removal rate of material in large volcanic clouds, a possible impact analog, does not occur by individual particle sedimentation, but rather by settling of agglomerates (Folch et al., 2010). Hence, removal in the region near the impact site may have been larger than Pope estimated, requiring a larger volume of quartz, or the removal of clastics

may be different than that of quartz. The value in Artemieva and Morgan (2009) for the pulverized material ejected from the crater is 3 orders of magnitude larger than the estimate of Pope (2002). Most of this material is in the ejecta curtain, not in the impact fireball, and so is deposited close to the impact crater. The shocked quartz is primarily associated with the impact fireball, so the bulk of the pulverized material may not be seen in Pope's analysis.

Pope assumed that quartz composed 50 % of all the clastic debris, so that all of the clastics injected weighed about 10^{16} g. This number is about 2 orders of magnitude less than the clastics from the Toba eruption (Matthews et al., 2012), and more than 3 orders of magnitude less than the Artemieva and Morgan (2009) estimate for clastics from the Chicxulub impact.

The assumption by Pope (2002) that quartz is 50 % of all the clastics is likely in error. There is no reason to think there is much quartz in the upper layers of sediment at the Chicxulub site. In the stratigraphic columns shown by Ward et al. (1995) the pre-impact sediments at Chicxulub consist of approximately 3 km of Mesozoic carbonates and evaporites with $\sim 3\text{--}4\%$ shale and sandstone. Therefore, it is more likely the quartz originates from the basement rocks. There is also not a strong connection between the physical processes that distributed the quartz (the impact fireball with high ejection velocity), and those that distributed the pulverized material (the ejecta curtain with low velocity).

It is possible that the quartz to clastics ratio is determined by the ratio of quartz to total debris in the samples closest to Chicxulub, since these may have suffered the least removal by sedimentation. Pope suggests these intermediate distance layers contain about 1 % quartz, but only considers the fireball layer, which is less than 10 % of the total ejecta layer within 1000 km of the crater. The remainder of the intermediate distance layer contains little quartz, so the clastics could be more than 1000 times the mass of the quartz. It is not clear that 1000 is an upper limit to the ratio of clastics to quartz because the quartz and pulverized material move along different paths in the debris cloud. If we accept this ratio of 1000 for the ratio of clastics to quartz, the mass of clastics from Pope's analysis would be 5×10^{18} g, which is within a factor of 6 of the Artemieva and Morgan (2009) value. If 1 % of this mass is submicron then 5×10^{16} g of submicron clastics would have been injected into the upper atmosphere.

Reconciliation of Pope (2002) and Toon et al. (1997) estimates of submicron clastics

Table 5 shows that the new estimate of submicron mass following the procedure of Toon et al. (1997) agrees with the new estimate following the procedure of Pope (2002) within 20 %. The new estimate is about 12 times less than the Toon et al. (1997) value mainly because Toon et al. (1997) did not consider that most of the pulverized mass would not be ejected from the crater. The new application of the

Pope (2002) approach leads to estimated submicron dust emissions that are about 500 times larger than the one originally derived by Pope (2002). The major difference is that we have assumed the ratio of quartz to clastics is about 1000, rather than 1 as assumed by Pope (2002). Despite the perhaps coincidental agreement of these two estimates, there is substantial uncertainty in the true mass of submicron clastic particles in the K-Pg distal layer. Observations of the submicron material in the distal layer are needed.

2.4.2 Submicron pulverized rock from a 1 km diameter impactor

In order to determine the properties of the pulverized ejecta from a 1 km impactor, we use the pulverized mass injection per teragram of impact energy from Toon et al. (1997), but reduce it by the factor of 25 discussed earlier to account for the fraction of the clastic mass with enough velocity to escape the crater. This procedure yields a clastic mass of 1.3×10^{16} g. For reference, the volume of clastics from the eruption of Mt. Tambora in 1815 is estimated to have been about 150 km^3 , which is a mass of about 3×10^{17} g. Hence, the Tambora eruption likely surpasses the clastics from the hypothetical 1 km diameter impactor by more than a factor of 10. The same size distribution for the clastics is recommended for the 1 km impact and the Chicxulub impact, since it seems to hold for a range of volcanic events from Mt. St. Helens to Toba, which span the 1 km diameter impactor in terms of clastics. We also suggest that the mass be initially mixed uniformly in the vertical above the tropopause. According to Stothers (1984), the Tambora clastics were deposited in layers that are centimeters in thickness at distances 500 km from the volcano. Accounting for the drift of the ash downwind, the area of significant ash fall was about $4.5 \times 10^5 \text{ km}^2$. If this same area is used for the initial injection of the clastics for the 1 km impact, then the column mass concentration is about 8.7 g cm^{-2} , which in turn is slightly less than 1 % of the atmospheric column mass. The estimated optical depth of the clastics in Table 3 is about 25 % of the optical depth from nanoparticles originating from vaporized rock. Given that these materials are much less absorbing than soot, and lower in optical depth than nanoparticles, they can probably be neglected in estimates of the climate changes due to a 1 km diameter impact on land.

3 Gas injections

There are a large number of gases that might be injected into the atmosphere after an impact and might be important to atmospheric chemistry, climate, or both. These can originate from the impactor itself, from ocean or groundwater, or from the target sediments. They may also originate in response to environmental perturbations, such as wildfires,

or atmospheric heating from the impact fireball and ejecta. Various estimates have been made for each of these sources. However, clear evidence from the distal layer is not available for any gases of potential interest. Some gases, such as carbon dioxide, would have stayed in the gas phase rather than condensing into particulate form. Other gases, such as those containing sulfur, may have reacted on the particles composing the distal layer, or formed independent particles. In either case, sulfur is so common in the environment it is difficult to detect an injection. For these reasons, all the gas-phase injections are uncertain. Below, we first discuss the chemical content of each of the potential sources of gases, and then we discuss the likely amounts of each material injected following an impact. Relevant ambient abundances are given in Tables 2 and 4 along with estimated injections for the Chicxulub impact and a 1 km impact. The ambient masses are given to assist the reader in understanding the magnitudes of the injections. Generally ambient concentrations are given in the literature in terms of the mixing ratio. To compute the masses we assume the ambient mixing ratios are constant over the whole atmosphere, or the stratosphere. We then convert the volume mixing ratio to the mass mixing ratio using the molecular weight and then multiply by the mass of the atmosphere above either the surface or tropopause to obtain the total mass of the gas. The ambient abundances assume the current stratospheric mixing ratio of Cl is 3.7 ppbv (Nassar et al., 2006), Br is 21.5 pptv (Dorf et al., 2006), inorganic I is 0.1 pptv (upper limit from Bosch et al., 2003), CO_2 is about 395 ppmv, and methane is about 1.8 ppbv. Stratospheric S, taken from the Pinatubo volcanic eruption, is about 10 Tg (Guo et al., 2004); reactive nitrogen, NO_x , in the stratosphere is difficult to quantify simply. Instead, we compare with the ambient abundance of N_2O in the stratosphere, about 2×10^{14} g N. N_2O is a major source of NO_x .

3.1 Impactor

3.1.1 Composition of the impactor

Kring et al. (1996) summarized the S, C, and water contents of a large number of types of asteroids. Trinquier et al. (2006) found from chromium isotopes that the Chicxulub impactor was most likely a carbonaceous chondrite of CM2 type. Such asteroids have 3.1 wt % S, 1.98 wt % C, 11.9 wt % water, and a density of 2.71 g cm^{-3} . Over the range of chondrites, which constitute 85% of meteorite falls, S varies from 1.57 to 5.67 wt %, C from 0.04 to 3.2 wt %, and water from 0.2 to 16.9 wt %. Kallemeyn and Wasson (1981) report that by mass CM carbonaceous chondrites contain about 4 ppm Br. Goles et al. (1967) report that Cl ranges from 190–840 ppm of carbonaceous chondrites, Br ranges from 0.25 to 5.1 ppm, and iodine ranges from 170 to 480 ppb. Table 6 summarizes the composition of asteroids using values for CM2 type carbonaceous chondrites from Kring et al. (1996) for S, C,

Table 6. Impactor composition, seawater composition, Yucatan impact site composition, and forest fire emission ratios.

	S	C	H ₂ O	Cl	Br	I	EC	N
Carbonaceous chondrite (g g ⁻¹ impactor)	3.1×10^{-2}	1.98×10^{-2}	11.9×10^{-2}	4.7×10^{-6}	3.27×10^{-6}	4.8×10^{-7}		
Seawater (g g ⁻¹ seawater)	9.1×10^{-4}	3×10^{-6}	0.965	1.9×10^{-2}	8.2×10^{-5}	6.0×10^{-10}	–	–
Impact site (g g ⁻¹ site)	7.1×10^{-2}	9.6×10^{-2}	0.07					
Emission ratios for forest fires g g ⁻¹ of dry biomass burned	2.9×10^{-4a}	4.3×10^{-1} as CO ₂ 4.4×10^{-2} as CO 5.1×10^{-3} as CH ₄	Highly variable, can equal dry weight	As CH ₃ Cl 1.4×10^{-5} to 1.3×10^{-4}	As CH ₃ Br 6.7×10^{-6}	As CH ₃ I 6.1×10^{-6}	6.6×10^{-4b}	7.5×10^{-4} as NO 6×10^{-5} as N ₂ O

^a The mass is given in terms of S, but the emission is in the form of SO₂. ^b We used 0.03 g g⁻¹ in Table 3, because forest fires will not produce as much soot as mass fires.

and water, and for the Mighei (the CM2 type example) from Goles et al. (1967) for Cl, Br, and I.

3.1.2 Gases from the impactor

Tables 2 and 4 indicate the direct contributions from 1 and 10 km impactors of a number of chemicals, as discussed further below. We assume that the entire 10 or 1 km diameter impactor melted or vaporized so that all of the gases are released. For the 10 km impactor, these gases would have been distributed globally in the hot plume along with the melt spherules within hours. They would reenter with the same vertical distribution as the Type 2 spherules. For the 1 km diameter impactor, the initial injection may have only covered half the Earth, with global distribution over days via wind, after reentry into the upper atmosphere.

We further assume that the vapors under consideration do not react with the hot mineral grains either in the plume or in the hot layer at the reentry site. In fact, given the large particle surface areas in the atmosphere over the globe it is possible that there was a significant transfer of material from the gas phase to the surfaces of the mineral grains in a short period of time.

As pointed out by Kring et al. (1996) and Toon et al. (1997), the S in a 10 km diameter impactor would exceed that from the Mt. Pinatubo volcanic injection by a factor above 1000. Even a 1 km diameter carbonaceous chondrite could deliver several times as much sulfur to the atmosphere as the Mt. Pinatubo eruption in 1991. Stratospheric water could be enhanced by a factor of more than 100 from the water in a 10 km impactor. Cl could be enhanced by factors above 500, Br by almost 500, and I by more than 50 000. However, there is not enough C in a 10 km asteroid to affect the global carbon cycle significantly.

Many investigators have pointed to sulfate as an important aerosol following the Chicxulub impact. Tables 1 and 3 compare the mass of sulfur from the impactor with the mass of the spherules and nanoparticles. The optical depth, which controls the climate change following the impact, and the particle surface area, which likely controls chemistry, are approximately linear with the mass. In our estimates, the sulfate coming directly from the asteroid could have a large optical depth assuming it was not removed on the spherules or large clastics.

3.2 Seawater

3.2.1 Composition and depth of seawater

The composition of seawater is given in Table 6 (Millero et al., 2008). It is thought that injections of water into the upper atmosphere will lead to droplet evaporation, with small crystals of salt left behind. If liquid water is left after a massive injection of water, the droplets will likely freeze leaving salt behind as particles embedded in ice crystals. Vaporization of water during the impact may leave behind salt crystals, or the salts may decompose into their components. As discussed by Birks et al. (2007), complex simulations are needed to determine how much material is freed from the salt particles to enter the gas phase where it might destroy ozone. In Tables 2 and 4, we list the total amounts of several interesting chemicals that might be inserted into the stratosphere. However, all of them except water vapor are likely to be in the form of a particulate until photochemical reactions liberate them.

A significant uncertainty related to any oceanic contribution to atmospheric composition is the depth of the ocean in relation to the size of the impactor, and the water content of sediments at the crater site. The depth of the ocean at Chicxulub at the time of the impact is not known. Many investigators have referred to it as a shallow sea. However,

Gulick et al. (2008) estimates that the water depth averaged over the impact site was 650 m, which is considerably deeper than earlier estimates. We use a water depth of 650 m in Table 2 to estimate the amounts of material injected by Chicxulub. A 1 km diameter impactor is smaller than the average depth of the world oceans, which is about 3.7 km.

3.2.2 Gases from seawater: Chicxulub

For the Chicxulub impact, we follow Pope et al. (1997) and assume that the 650 m depth of seawater within the diameter of the impactor (10 km) will be vaporized, follow the path of the Type 2 spherules, and reenter the atmosphere globally. In Table 2 we compute the water vaporized following the equations in Toon et al. (1997). These equations, assuming an impact velocity of 20 km s^{-1} , led to an order of magnitude greater injection of water than using Pope's estimate. The vaporized water is 0.4 times the impactor mass. During the vaporization of the seawater we assume the water will be present as water vapor, and that the materials in the water will be released as vapors. Some of these materials likely would react quickly with the hot minerals in the fireball or later with the hot minerals in the reentry layer.

It is also likely that a considerable amount of water was splashed into the upper atmosphere. Ahrens and O'Keefe (1983) estimated that the water splashed above the tropopause from a 10 km diameter impact into a 5 km deep ocean would be 30 times the mass of the impactor. We assume that the amount of water splashed above the tropopause will scale linearly with the depth of the ocean. Therefore, about 4 times the impactor mass of water may have been splashed into the upper atmosphere. Much of this water may immediately condense and rainout, as discussed in Toon et al. (1997). However, some of the dissolved salts may be released if some of the water evaporates. The assumed injection of gases, and particulates that might become gases, from the ocean is summarized in Table 2 for the Chicxulub impact.

3.2.3 Gases from seawater: 1 km asteroid

No seawater is injected by the 1 km diameter asteroid impact on land. If a comet hit the land there would be a water injection.

Pierazzo et al. (2010) estimated that 43 Tg of water would be injected above 15 km by a 1 km asteroid impact into the deep ocean. Of this water, 25 % is in the form of vapor and 75 % in the form of liquid water. In their modeling the water was assumed to be distributed with a uniform mixing ratio from the tropopause to the model top. It was also spread uniformly over an area $6200 \times 6200 \text{ km}$ in latitude and longitude. Using the equations in Toon et al. (1997) for the vaporized water produces a value which is 60 % of the vaporized water from the detailed modeling used in Pierazzo et al. (2010). Given these water injections we use the composition of seawater to determine the injections of the various

species. Pierazzo et al. (2010) estimate injections of Cl and Br that are more than an order of magnitude smaller than ours because they consider the amounts that have been converted into gas-phase Cl and Br by photochemical reactions in the atmosphere, while we estimate the total injections, which initially are likely to be in the particulate phase.

3.3 Impact site

3.3.1 Composition of the impact site

The sea floor at the Chicxulub impact site, like the modern Yucatan, contained abundant carbonate and sulfate-rich deposits. Ward et al. (1995) conclude that 2.5–3 km of sedimentary rock were present at Chicxulub, composed of 35–40 % dolomite, 25–30 % limestone, 25–30 % anhydrite, and 3–4 % sandstone and shale. The dolomite and limestone are no doubt porous. Pope et al. (1997) estimate the carbonates in the Yucatan have a porosity of 20 %. The pores would have been filled by seawater since the sediments were submerged. This groundwater produces an equivalent water depth of about 400 m. The carbon content of limestone is 12 % by weight and of dolomite 15 % by weight. The sulfur content of anhydrite is 23.5 % by weight. To our knowledge, trace species such as Br, Cl, and I have not been reported for these sedimentary rocks, but would be present in the seawater in the pores.

3.3.2 Gases from the impact site

For the 10 km Chicxulub impact, we follow Pope et al. (1997) for the abundances of S and C assuming 30 % anhydrite, 30 % limestone, and 40 % dolomite. The composition of the impact site is given in Table 6. We ignored species other than S and C that might be in the target material. It is difficult to follow the target debris since some of it is vaporized, and some melted. We follow Pope et al. (1997) and assume that the upper 3 km of the target is vaporized within the diameter of the impactor. The gases within this volume of vaporized material are assumed to be released, and to follow the trajectories of the Type 2 spherules. Pope et al. (1997) estimated the amount of material that would be degassed from target material that was melted or crushed in a large impact. We use the values from Table 3 of Pope et al. (1997) for out of footprint vapors in our Table 2 for the degassed impact site emissions. We also assume that the granite underlying the impact site does not contribute.

The source gases from a 1 km land impact would depend on the composition of the impact site, so we do not list values in Table 4. We assume nothing would be liberated from the sea floor in a 1 km impact in the deep ocean.

3.4 Fires

3.4.1 Composition of smoke

It is well known that forest fires emit a wide variety of vapors into the atmosphere. Andreae and Merlet (2001) provide emission ratios (gram of material emitted per gram of dry biomass burned) for many vapors expected to be important in the atmosphere, as listed in Table 6. As discussed in Sect. 2.2.1, the soot emission may have been enhanced relative to wildfire estimates by Andreae and Merlet (2001) after the Chicxulub impact because the impact-generated fires were mass fires. We do not consider any enhancements of the gas-phase emission ratios, but they may also be impacted by fire intensity or the types of plants making up the biomass.

3.4.2 Gases from fires

In Tables 2 and 4, we computed the burned mass from Chicxulub assuming that 1.5 g cm^{-2} of dry biomass burns over the entire land surface area of the Earth, and then used the emission factors from Andreae and Merlet (2001) to obtain the gas-phase emissions. For a 1 km impact, we assume the area burned is $4.1 \times 10^4 \text{ km}^2$ (Toon et al., 1997), and the dry biomass is 2.25 g C cm^{-2} . We then used the emission ratios from Andreae and Merlet (2001) to compute the gas-phase emissions. Comparing the gas-phase emissions from fires in Tables 2 and 4 with ambient values indicates that there would be large perturbations for all gases for the 10 km diameter impact. Only iodine is significantly perturbed for the 1 km impact. For the gas-phase emissions, we suggest using the same vertical profile as suggested for soot earlier. The emissions would only occur over the region near the impact site for the 1 km impact.

3.5 Gases generated by atmospheric heating

The energy deposited in the upper atmosphere by the initial entry of the bolide, as well as by the rising fireball, may have converted some N_2 to NO_x . Early studies suggested that a large fraction of the impact energy would be put into the lower atmosphere, which in turn led to suggestions that a large amount of nitrogen oxides would be produced from the heated air. However, it is now understood that most of the energy release from an impact to the atmosphere will occur at high altitude from reentry of spherules and other debris. Toon et al. (1997) reviewed the various ways in which NO_x might be generated following an impact, largely following Zahnle (1990). They concluded that $3 \times 10^{16} \text{ g}$ of NO might be produced from the atmosphere for a 10 km diameter impact with about half coming from the plume at the impact site, and half from the reentry of material across the Earth. We have recorded this value in Table 2. For comparison, Parkos et al. (2015) conducted detailed evaluations of the NO_x produced by the infalling spherules and concluded the spherules could produce 1.5×10^{14} moles of NO_x

($3 \times 10^{15} \text{ g}$ if the NO_x is in the form of NO) which they further concluded was not sufficient to acidify ocean surface waters. In Table 2 we use the Toon et al. (1997) injection of NO since it includes both source mechanisms. According to Zahnle (1990), a 1 km impact on land might produce $0.6 \times 10^{14} \text{ g}$ of NO, largely in the hot plume at the impact site. This value is entered in Table 4. For comparison, we note that Pierazzo et al. (2010) suggested that the mass of NO produced by a 1 km ocean impact is about $0.39 \times 10^{14} \text{ g}$.

3.6 Discussion of gas injections

Some of the gas-phase sources just discussed are easy to apply to an impact. For example, the emissions from fires simply depend on the area burned, the fuel loading, and the emission factors.

Other sources of gases are more difficult to evaluate. Since we have no measurements for large impacts, the form of emission can be uncertain. For example, sulfur could be injected as SO_2 or SO_3 . Another difficulty that comes in understanding the contribution of target material to gases such as SO_2 is the pressure needed to vaporize the material. Pope et al. (1997), for example, adopted pressures above 70 GPa to vaporize carbonate, 100 GPa for complete vaporization of anhydrite, and 10 GPa for water vaporization from pores. These vaporization pressures are higher than suggested by early researchers, leading to lower amounts of target vaporized. Pierazzo et al. (2003) redid the impact calculations and also estimated the amounts of materials that might be released, which are close to those estimated by Pope et al. (1997). The altitude distribution of the ejecta varies with the source of the material. Finally the chemical form of the emission varies with thermochemistry in the ejecta plume or fireball, and interactions with hot mineral surfaces, and for some materials exposure to high temperature on reentry.

Tables 2 and 4 summarize our choices for the injections of the various gases. For each type of source, we also specify the altitude of the expected injection, using a reference to Tables 1 and 2 for the particle injections. We assume all of the impactor mass entered the rising fireball, so it would be injected near 60 km altitude along with the spherules. In some cases, for example for the degassed target material and for splashed seawater, we consider the material to have been uniformly mixed above the tropopause. For materials coming from fires we assume the same vertical injection as for soot.

As has been pointed out many times (Kring et al., 1996; Toon et al., 1997; Pope et al., 1997; Pierazzo et al., 2003) the sulfur injection from a 10 km impactor might be thousands of times greater than that from the Pinatubo eruption, and also was likely larger than the injection from the massive Toba eruption by a factor between 10 and 100. Our sulfur injection from the target material is about half that of the Pope et al. (1997) estimate of 10^{17} g and slightly less than the Pierazzo et al. (2003) estimate for a 15 km diameter impactor of $7.6 \times 10^{16} \text{ g}$. Our sulfur injection from the asteroid

itself is within the range suggested by Pope et al. (1997) of $2.7\text{--}5.9 \times 10^{16}$ g. Interestingly, the sulfur injection we estimate for Chicxulub is about 10 times greater than the yearly emission estimated by Schmidt et al. (2016) for a large flood basalt from the Deccan traps. Of course, the flood basalt might continue for a decade or more, bringing the total sulfur emission close to that from the Chicxulub impact. Table 4 suggests that the sulfur injection from a 1 km impact would be several times greater than that from the Pinatubo eruption, but that would be only a modest injection relative to historical volcanic eruptions. In Tables 1 and 3 we assume the injected sulfur gas is converted into sulfate. If so, it would yield a large optical depth for the Chicxulub impact. However, for both the 1 km and Chicxulub impacts, the sulfur injection, if converted to sulfate, would be an order or magnitude less massive than the nanoparticles. Therefore, the sulfate would be an order of magnitude less important optically than the nanoparticles. While it might exceed the soot mass slightly, soot is much more important optically than sulfate, which is transparent at visible wavelengths. Therefore, the sulfate in our model is of relatively little importance optically, unless the sulfur remains in the air after the other particles are removed.

Our estimated C injection (in the form of CO_2) is dominated by emissions from forest fires. We have the same emission from the impactor as Pope et al. (1997), but we have less than half the emission from the target material as Pope et al. (1997) or Pierazzo et al. (2003). All these studies suggest a small impact perturbation relative to the CO_2 65 million years ago, which was several times larger than now.

The water vapor injections in Tables 2 and 4 are very large compared with ambient values in the stratosphere. However, most of the water is from fires, and half will be injected into the troposphere where it will be quickly removed. The water from the impactor and target is modest, about 1 cm as a global average depth of rain. The typical rainfall averaged over the current Earth is about 3 mm day^{-1} . The emissions from the impactor and from vaporized seawater, both of which would have been injected globally at the same altitudes as the Type 2 spherules, are capable of saturating the entire ambient stratosphere. Our water injection is similar to that estimated by Pope et al. (1997) and Pierazzo et al. (2003). While the water vapor has been largely ignored in previous work on the Chicxulub impact, it has the ability to alter the thermal balance of the stratosphere by emitting and absorbing infrared light. Water vapor may have been a factor in the radiation of thermal energy to the surface during the first few hours after the K-Pg impact, since Goldin and Melosh (2009) sought an infrared absorber to prevent radiation from escaping from the top of the atmosphere. Some of the particles in the stratosphere might be removed by precipitation, but the mass of water injected is comparable to the mass of the nanoparticles and spherules. Therefore, removal by precipitation is probably not significant since if the water condenses on all the particles it will add only a small mass,

and increase the fall rate only slightly, while if water condenses on only a subset of the particles it will remove only a subset. The water injection by the 1 km diameter impact on land is about 15 % of the ambient water, but might still lead to some significant perturbations if it is injected into the upper stratosphere. The 1 km impact in the deep ocean could inject about 40 times the ambient water into the stratosphere (Pierazzo et al., 2010), and water should be considered in simulations of such impacts.

For the 10 km diameter impactor, there are injections of Cl, Br, and I that exceed the ambient values by orders of magnitude. There are significant sources for all three halogens from fires, the impactor and seawater, so it seems inescapable that large injections would have occurred. The injections of NO_x from fires, and from heating the atmosphere are also very large compared with ambient values. For instance, Table 2 shows the NO_x injections are 1–2 orders of magnitude larger than the stratospheric burden of N_2O , the principle source of NO_x . For the 1 km diameter land impact, only the injections of I and NO_x appear large enough to perturb the chemistry of the stratosphere. However, as discussed by Pierazzo et al. (2010) significant Cl and Br injections could occur for a 1 km impact in the ocean. Seawater injections of Cl, Br, I, and S are complicated because the salts may be injected in particulate form.

4 Implications for climate, atmospheric chemistry and numerical modeling, and suggestions for future data analysis

Since the discovery of the K-Pg impact by Alvarez et al. (1980), many papers have speculated on which of the many possible effects of the impact on the environment could have caused the mass extinction. It has become fashionable to claim that one or another effect is dominant. However, it is quite likely that several effects overlapped, each of which might have been devastating to a particular species or ecosystem, but which together made survival very difficult for a broad range of species distributed over the globe. Here, we summarize the environmental perturbations we find likely. However, there are many uncertainties, and additional data are needed. We outline the data that would be useful to obtain from the geologic record, and summarize them in Table 7. Also, models have barely scratched the surface of what is possible in better understanding of the post-impact environment. We summarize the types of modeling work that would be interesting to pursue. We extend these ideas to smaller impacts since more than 50 impacts of kilometer-sized objects may have occurred since the extinction of the dinosaurs.

Table 1, shows that spherules, soot, nanoparticles, submicron clastics, and sulfates each may have had very large optical depths. An optical depth greater than unity could have serious consequences for the environment if maintained for

Table 7. Suggestions for data collection.

Property of interest	Rationale
Global distribution of spherules	Some impact models suggest spherules were not distributed globally, limiting area of Earth that might experience fire ignition
Number concentration, size of spherules	Current data are incomplete on number and size of spherules
Soot distribution	Profile soot/iridium/spherule distribution to determine if fires are contemporaneous with iridium fallout
Nanometer material	Nanometer material has been detected, but its mass needs to be quantified
Clastics	Submicron component not detected; possibly search for micron/submicron shocked quartz
Sulfur	Use sulfur isotopes to search for extraterrestrial sulfur, sulfur mass-independent fractionation (MIF) to test for prolonged lifetime

very long. Each of these materials was likely present in the atmosphere, so they may have interacted.

The spherules are unlikely to have changed climate directly because they would have been removed quickly from the atmosphere by sedimentation due to their large size. However, these particles, together with the other impact debris with significant mass, likely heated the upper atmosphere to temperatures between 1000 and 2000 K. The high-temperature upper atmosphere would then have irradiated the surface with near-infrared radiation, causing forest fires. Wolbach et al. (1985) first recognized that the global biota likely burned after the impact, and Melosh et al. (1990) identified the mechanism for starting the fires. The recent work by Goldin and Melosh (2009) identified some complexities in the ignition mechanisms that need further work to be understood. They pointed out that the light might be blocked by the large spherules falling below the heated atmospheric layer. However, this is a complex problem since water vapor and the vaporized impactor would have been present to block radiation escaping to space. Also convection should occur in such a strongly heated layer, which would act to retard the fall of the particles as it does for hailstones in tropospheric convection. Moreover, the mass of debris injected at 70 km, as assumed by Goldin and Melosh (2009), greatly exceeds the mass of air. This mass distribution is unstable and would lead to rapid stirring of the atmosphere down to 50 km. These issues all deserve further study with suitable models. Furthermore, evidence for the nanoparticles should be sought as discussed further below.

Robertson et al. (2004) argued that large dinosaurs and other unsheltered animals could have been killed immediately by the radiation from the sky and the subsequent fires. However, it is possible there were refugia on the land, either in regions where spherules did not reenter the atmosphere, as suggested by Kring and Durda (2002) as well as Morgan et al. (2013), or in regions that happened to have heavy cloud cover which may have blocked the radiation. To better under-

stand the possibility of refugia, more complete evidence for the global distribution of spherules would help resolve their possible nonuniform deposition, as suggested in Table 7. It is known that iridium was perturbed worldwide following the K-Pg impact. Although iridium concentrations are spatially variable for a number of reasons, they are basically homogenous over the Earth and do not fall off with distance from the impact site, or at high latitudes. Similar data on spherules would be useful to determine if the spherules were injected everywhere or in special places. Numerical values of the spherule concentrations and size distributions to augment the values noted by Smit (1999) would also be of value, as noted in Table 7. Models of the transmission of the light from the hot debris layer above 60 km through dense water clouds and the response of the clouds to the heating would be also useful. It has long been recognized that intense thermal radiation and fires could not have been the only extinction mechanisms at work, since the mass extinctions in the oceans could not have occurred in this way, but instead were likely due to the low light levels preventing photosynthesis (Milne and McKay, 1982; Toon et al., 1982, 1997; Pollack et al., 1983; Robertson et al., 2013b). The low light levels would have been caused by the high optical depths of the soot and nanoparticles that remained suspended in the air for a year or more after the impact.

We know from the work of Wolbach et al. (1985, 1988, 1990a, b, 2003) that there is abundant soot in the K-Pg distal layer. It is highly likely that the soot originated from wildfires (Robertson et al., 2013a), but its origin is of secondary concern for climate. The widespread distribution of the soot in the layer, and the small size of the particles indicate this material was almost certainly global in extent. Wolbach et al. (1988, 1990b) show that soot and iridium are tightly correlated across the K-Pg distal layer. The soot and iridium in the distal layer must have been deposited within a few years of the impact, since small particles will not stay in the air much longer. Therefore, any fires must have been within a

year or two of the impact. As noted in Table 7, further examination of the distributions of soot, iridium and spherules might clarify how long these materials remained in the atmosphere, which is expected to be days for the spherules, and a few years for the soot and iridium on small particles. Once in the water column, spherules would fall to the bottom in days or weeks. However, in the absence of fecal pellets formed by plankton around the soot, it would take decades for soot to reach the ocean depths by falling. Currents would likely carry the soot down rather than gravity.

The amount of soot in the K-Pg distal layer would produce a very high optical depth when it was in the atmosphere. The transmission of light depends not only on the optical depth but also on the single-scattering albedo of the particles. The single-scattering albedo measures the fraction of the light that is scattered or absorbed. Scattering light, which occurs from sulfates that absorb sunlight only weakly, is not nearly as effective in changing climate as absorbing light.

As discussed by Toon et al. (1997), soot with an optical depth of 100 would prevent any sunlight from reaching the surface – it would be pitch black. No climate simulations of such large soot optical depths have ever been conducted. However, there have been simulations for optical depths in the range of 0.05–1, which show temperatures dropping to ice age conditions within days, precipitation falling to 50 % of normal, and the ozone layer being destroyed as discussed further below (Robock et al., 2007a, b; Mills et al., 2008, 2014). There are a number of complexities inherent in climate calculations for soot. For example, it is important to know how long the soot remained in the atmosphere in order to determine how long photosynthesis may have been retarded in the oceans. The lifetime of the soot in turn may depend on the size of the soot particles, their shape, the amount of rainfall in the lower atmosphere, and the amount of sunlight reaching the soot. The amount of sunlight reaching the soot matters because heating the soot also heats the surrounding air, causing it to rise and loft the soot to high altitudes, where it is protected from rainout (Malone et al., 1985; Robock et al., 2007a, b). These issues can be considered in modern climate models.

Much of the vaporized impactor and target material is thought to have recondensed to 250 μm sized spherules (O'Keefe and Ahrens, 1982; Johnson and Melosh, 2012b), which are observed, but a significant fraction may have remained as nanometer-sized grains (Johnson and Melosh, 2012b). Iron-rich nanophase material with a diameter of 15–25 nm has been identified in the fireball layer at a variety of sites by Wdowiak et al. (2001), Verma et al. (2002), Bhandari et al. (2002), Ferrow et al. (2011) and Vajda et al. (2015) among others. However, the abundance of this nanophase material is not yet constrained by observations. As noted in Table 7, it is important to quantify the abundance of this nanophase material and to confirm that it is the remnant of the vaporized target and impactor. If the amount of vapor remaining at the end of the Johnson and Melosh (2012b) cal-

ulation is roughly the amount that remained as rock vapor in the atmosphere, given the optical depth estimate in Table 1 and its input location in the upper atmosphere above the soot generated by forest fires, this nanophase material would be the dominant source of opacity for changing the climate, and would also greatly affect the amount of radiation emitted to the surface that could start wildfires in the hours following the impact. The material contains iron, so it is likely to have been a good absorber of sunlight. Alternatively, this material might have attached itself to the large spheres and been quickly removed, though this seems unlikely since the large spheres would separate gravitationally from the smaller material within hours. No one has yet considered the effect of this nanophase material, which is distinct from the clastics envisioned by Toon et al. (1997) and Pope (2002) on the environment after the K-Pg impact.

The most massive part of the ejecta from the K-Pg crater consisted of clastics: crushed and pulverized material. Much of this material fell relatively close to the crater, though significant amounts were emplaced as far as 4000 km from Chicxulub. For comparison, the Toba volcanic eruption about 70 000 years ago is estimated to have released more than 2×10^{18} g of clastics (Matthews et al., 2012), a factor of about 15 less than our estimate for the Chicxulub impact in Table 1, but more than 200 times greater than the upper limit previously estimate by Pope et al. (1997) for the clastics generated by Chicxulub.

The Toba eruption may have had a significant impact on the climate, as discussed further below; however, the magnitude of the effect is controversial. Alvarez et al. (1980), as well as Toon et al. (1982) and Pollack et al. (1983), thought that the K-Pg layer was dominated by submicron clastics that caused major loss of sunlight at the surface and consequently very low temperatures. However, while we do not know the fraction of the layer composed of submicron clastics, it is clear that the layer is both thinner than thought in the years just after its discovery and also dominated by other parts of the impact debris such as the spherules and the nanoparticles. It would be very useful to measure the amount of submicron clastics in the K-Pg distal layer. Possibly, as suggested in Table 7, one could start by identifying the amount of submicron quartz in the layer by searching for small shocked quartz grains. Toon et al. (1997) and Pope (2002) used two differing indirect approaches to quantify the submicron clastics, and came up with answers that differ by a factor of about 10^4 . Here, we attempted to reconcile these approaches, with the result shown in Table 1 yielding a significant optical depth. Although the submicron clastics by themselves would have produced extreme climate changes if they were as abundant as we estimate, they would have been less important than the soot, and the nanoparticles given our estimates here. The submicron clastics may have been injected higher than the soot, but lower than the nanoparticles on average. Climate calculations involving all these materials are needed to understand how they may have interacted in the atmosphere.

The final particulates with large optical depths in Table 1 are sulfates. Pope et al. (1997), Pierazzo et al. (2003), and others have advocated for the importance of these particles in recent years. Unfortunately, sulfates in the K-Pg layer have not been traced unambiguously to the impact, because sulfur is so common in the environment. Possibly sulfur isotopic studies could distinguish the sulfur in the impactor from sulfur in the terrestrial environment, but we are not aware of such studies. While there is little doubt that large amounts of sulfur were present in the target material and in the asteroid, it is possible that much of it reacted with the hot rock in the impact plume, or the atmospheric layer heated by re-entering material. Sulfur is present in impact melt spherules and in carbonaceous clastics, so not all of it was released to the gas phase. Given the large opacity of the numerous types of particles in the atmosphere, photochemical reactions would have been inhibited, which would retard the conversion of sulfur dioxide gas into sulfate particles. It is possible that measurements of the sulfur MIF could reveal whether the sulfur quickly reacted with rocks, which should yield a MIF of zero, or if the sulfur slowly converted to sulfate, which might lead to MIF not being zero if resolved over the thickness of the distal layer. It is known that a non-zero MIF can occur following volcanic eruptions due to time-dependent movement of sulfur between changing sulfur reservoirs in the atmosphere (e.g., Pavlov et al., 2005).

It is not clear if SO_3 or SO_2 was the dominant sulfur bearing gas in the ejecta plume. However, the gas-phase reaction of SO_3 and water is not a simple reaction as often abbreviated in papers about atmospheric sulfur chemistry, but instead involves water vapor clusters or SO_3 adducts. Sulfur dioxide is observed to convert to particulates with an *e*-folding time of less than 1 month for moderate-sized volcanic eruptions such as the Mt. Pinatubo eruption. Following the K-Pg impact, sulfur dioxide or trioxide gas may have had an extended lifetime in the atmosphere, due to the lack of sunlight to drive chemical reactions to convert it to sulfates. Clastics and nanoparticles and soot may have coagulated to large sizes and fallen out over a year or two. Alternatively, the sulfur gases may have reacted quickly on all the surfaces present, particularly in hot water present in the hot radiating layer when the ejecta reentered. Pope et al. (1997) and Pierazzo et al. (2003) have pointed out the possible importance of the extended lifetime of the sulfate for causing a prolonged period without photosynthesis in the oceans. However, clastics or soot need to be present in the sulfate to achieve the loss of sunlight. Recent work on the Toba eruption (Timmreck et al., 2010) shows that large sulfur injections do not produce proportionately larger climate perturbations because the climate effects of sulfur injections are self-limiting, as originally shown by Pinto et al. (1989) and recognized by Pope et al. (1997) and Pierazzo et al. (2003). Toba probably injected an amount of sulfur dioxide within an order of magnitude of that from the K-Pg impact. Larger particles have smaller optical depths, and shorter lifetimes, than smaller particles that

result from smaller SO_2 injections. Further work is needed to understand the chemistry of the sulfur injected by the Chicxulub impact to determine if it was a significant factor in the extinction event.

Table 2 shows that significant injections of various ozone destroying chemicals such as NO_x , Cl, Br, and I likely occurred. The effects of these gases need to be considered in calculations but, given the expected darkness, photochemistry may have ceased until the atmosphere cleared.

Table 3 suggests that the much smaller mass injections from the impact of a 1 km diameter asteroid on land may produce optical depths that may still be important. Climate models are needed to fully evaluate these perturbations. At first glance the injections seem small. For example, the sulfur injection is only about 4 times larger than that from the Pinatubo eruption. However, the soot injection is very large. Robock et al. (2007a) and Mills et al. (2014) examined smoke injections at the tropopause of about one-third the 1 km asteroid injection near the tropopause and found that the ozone layer was severely damaged, and low enough temperatures resulted in damaged crops for a decade after the injection. Table 4 also indicates significant injections of iodine, which may further damage the ozone layer.

About 50 1 km impacts might have occurred since the demise of the dinosaurs. Based on the fraction of Earth covered by water, about 35 of these would be expected to have hit the oceans, perhaps resulting in large ozone losses as discussed by Pierazzo et al. (2010). Each of the 15 impacts that occurred on land might have led to significant injections of nanoparticles. Paquay et al. (2008) recognized the osmium signature of two large impacts in the late Eocene, which produce the 100 km diameter craters at Popigai and Chesapeake Bay. The osmium indicates a substantial input of vaporized impactor to the atmosphere from collisions of asteroids larger than 1 km in diameter. Climate model simulations are needed to evaluate the climate changes that might have occurred. The effects could have been variable for a variety of reasons, including variability in the light-absorbing properties of rock from differing objects. To have injected significant amounts of smoke the impactor would need to hit a tropical forest, or at least a heavily forested region. About 26 % of the world is currently forested; about 6 % is in tropical rain forest. Forested area has greatly declined. Tropical rainforests might have covered as much as 20 % of the Earth until recently. Hence, about three 1 km objects might have hit a tropical rainforest and injected significant amounts of smoke since the K-Pg event.

In this work, we have established a set of initial conditions (Tables 1–4) that may be used for modeling the climate and air chemistry after the K-Pg impact, or the impact of a 1 km asteroid. Other authors have considered some of these initial conditions, but some, such as the nanoparticles from the vaporized impactor, have not been previously studied in the detail needed to fully evaluate their importance. Much more work is needed to obtain field data to further

constrain some of parameters, and to resolve remaining differences of opinion about some of the values. However, simulations using these initial conditions can now be conducted with modern models of climate and atmospheric chemistry, which should shed light on the environmental conditions at the K-Pg boundary and the dangers posed by future impacts. We recently completed such simulations using the Whole Atmosphere Community Climate Model (WACCM) at the National Center for Atmospheric Research in a configuration similar to that used by Bardeen et al. (2008) and Mills et al. (2014).

5 Data availability

This paper uses data from the literature, which is referenced in the paper, and summarized in Tables 1–6.

Author contributions. Owen B. Toon worked to compile the particle and gas emissions. Charles Bardeen tested them in a climate model to determine if the initial conditions were specified completely. Rolando Garcia considered the gases that would be important for atmospheric chemistry.

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