Our finding that MACs serve as a direct substrate for methanogens may not be limited to coal-bed environments. In the deep subsurface, MACs are contained in sedimentary organic matter derived from lignin in higher plants, namely kergen, with quantitative variation depending on the maturity (24). In fact, alkyl-methoxyphenols with a short C4–C6 chain have been detected in the pyrolysates of immature kerogen (25). Kergen is ubiquitous in sediments and accounts for most of the organic matter in subsurface environments (26). Microorganisms from the genus Methermicoccus and related clones have often been detected in deep subsurface environments worldwide (fig. S5). Methoxytrophic methanogenesis may therefore play an important role in the biogeochemical carbon cycle in Earth as well as in the formation of biogenic gas, which accounts for more than 20% of natural gas resources, including CBM (27).

REFERENCES AND NOTES


ACKNOWLEDGMENTS

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SUPPLEMENTARY MATERIALS

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Materials and Methods

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GLOBAL CLIMATE CHANGE

Impact ejecta at the Paleocene-Eocene boundary

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Extraterrestrial impacts have left a substantial imprint on the climate and evolutionary history of Earth. A rapid carbon cycle perturbation and global warming event about 56 million years ago at the Paleocene-Eocene (P-E) boundary (the Paleocene-Eocene Thermal Maximum) was accompanied by rapid expansions of mammals and terrestrial plants and extinctions of deep-sea benthic organisms. Here, we report the discovery of silicate glass spherules in a discrete stratigraphic layer from three marine P-E boundary sections on the Atlantic margin. Distinct characteristics identify the spherules as microtektites and microkrystites, indicating that an extraterrestrial impact occurred during the carbon isotope excursion at the P-E boundary.

Ejecta from known impact strewn fields have common features that provide useful criteria for identifying previously undiscovered impact ejecta in the geologic record. An ejecta deposit typically results from an airfall event and occurs in a discrete stratigraphic layer. If spherules are present, they generally are glassy (microtektites) and may contain crystallites (microkrystites), both of which have characteristic morphologies and may be either solidified melt ejecta or vapor condensates (1–3). Microkrystites show distinct internal crystallite textures that indicate rapid quenching from high temperature and, if they formed as a vapor condensate, they can be enriched in projectile material (3). Microtektites that form as melt droplets solidify at slightly lower temperatures and are more likely to be vesicular; they may contain inclusions of high-temperature components such as lechatelierite, a quartz glass that solidifies above 1750°C (4, 5). Shock-metamorphosed grains can be associated with microtektites but are not uniformly distributed throughout a strewn field. For example, the presence and abundance of shocked grains is spatially heterogeneous in the widespread Australasian strewn field (for which an impact crater has yet to be identified, despite being the largest and youngest Cenozoic tektite event at 0.8 million years ago (Ma)). The Ivory Coast strewn field (next youngest at 1.1 Ma) that emanated from Bosumtwi crater contains (micro)tektites but no shocked mineral grains (see (6) for review).

We examined three marine Paleocene-Eocene (P-E) boundary sections that encompass the onset of the carbon isotope excursion (CIE) that defines the P-E boundary (Fig. 1): continental shelf sites Wilson Lake B (39.6598°N, 75.0472°W) and Millville [Ocean Drilling Program (ODP) Leg 174AX; 39.4045°N, 75.0889°W (6)], where the onset of the CIE coincides with the base of the thick Marlboro Clay unit of the Salisbury Embayment, and pelagic sediments at ODP Hole 1015B at Blake Nose (30.6253°N, 76.3758°W, water depth 1980.6 m) (9). To constrain the stratigraphic level of the CIE onset, we use detailed bulk sediment carbonate isotope records from Wilson Lake B and Millville (10) and Hole 1015B (12). We also report preliminary data from an exposure of the Marlboro Clay near Medford, New Jersey, to eliminate the possibility of spurious drilling-induced contamination as the source for the spherules [see (22)].

At the Wilson Lake B and Millville shelf localities, as well as Hole 1015B, peak spherule abundance occurs close to the P-E boundary as constrained by the δ13C decrease in bulk sediment (Fig. 1). Spherules are found over a ∼20-cm interval in each case, with the number of spherules increasing sharply up-section from zero below to a peak at or close to the P-E boundary. Maximum abundances in the >63-μm size fraction are between 2.5 and 3 spherules per g at Wilson Lake B and Millville and 4 spherules per g at Hole 1015B, followed by a gradual decline in abundance to zero over ~10 cm. At Wilson Lake B and Millville, the peak spherule abundance coincides with the

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basal contact of the Marlboro Clay, which is gradational with some reworking of the underlying Vincentown Formation facies.

These abundances are similar to impact spherule abundances in distal ejecta layers linked to other extraterrestrial impact events, such as the Late Eocene clinopyroxene (cpx)–bearing spherule layer associated with the Popigai impact crater (J3–15). Late Eocene cpx spherules that are found in ODP Site 738 in the southern Indian Ocean (~2 spherules per g (>150 μm)), in ODP Site 689 in the Southern Ocean (~5 to 6 spherules per g (>63 μm)) (see J3 for review), and at Massignano, Italy (~4 spherules per g (>150 μm)) (J6) show concentrations similar to ours.

The glassy spherules from Wilson Lake B and Millville range from 65 to 500 μm in diameter (average 302 μm), whereas those at Hole 1051B average 274 μm in diameter. They have generally spherical and rotational morphologies, as well as other characteristics of splash-form microtektites (J1, J7) (Fig. 2), and are occasionally colorless, translucent green, or dark-brown/black. The spherules often have surface pits (Fig. 2, A and C) and in some cases microcraters (Fig. 2G and fig. S7), indicating relative velocities high enough to fracture the spherules on impact with one another or other objects after solidification. Such microcraters would be an unexpected feature of volcanic spherules. There are also occasional dumbbell forms and smaller spherules accreted to larger ones (Fig. 2F), often of different chemical composition (Fig. 3E), suggesting that they were sintered to one another in a molten form and quenched rapidly. We conclude that the stratigraphic distribution and morphologies of the spherules are wholly consistent with the expected manifestation of an air-fall sedimentary event such as an impact ejecta bed, modified by minor and typical postdepositional bioturbation of marine sediments.

Major oxide chemistries of the P-E spherules show a wide range of compositions distributed equally among Wilson Lake B, Millville, and Hole 1051B (Fig. 4) (J2). The spherules are ~33 to ~50% SiO2 and have CaO contents between ~20 to ~35%, which is consistent with a CaCO3-rich target rock, with Al2O3 and FeO as the second and third most abundant oxides, at around 15% and 10% on average, respectively. This variability is typical of tektite spherules and consistent with an impact origin of the spherules (J8, J9). Impact melts have little time for homogenization, but a population of major oxide chemistries will generally follow broadly linear compositional trends with SiO2 content because they were generated from the same source rock at the site of impact (J8). In contrast, ashes or spherules from a volcanic eruption tend to exhibit inter- and intra-spherule compositional homogeneity (J20). Magma is generally well mixed during a single eruptive event, and hence volcanic spherules from the same eruption sequence will be compositionally homogeneous and generally have much higher water content than impact spherules [up to a few weight % (wt %)] (20]). The water content of impact glasses is generally much lower than volcanic spherules (21, 22) because water is removed by vapor stripping during solidification (23). Field transmission infrared spectroscopy (FTIR) on representative P-E spherules reveals water contents below 0.023 wt % (see table S5), much lower than expected for volcanic spherules but typical of impact-generated microtektites.

In cross section, the internal texture of most of the P-E spherules is glassy and relatively featureless (Fig. 3); exceptions are the less common microtektites (discussed below). The glassy spherules also occasionally have microcrysts just inside the rims and are mildly vesicular. This vesicular nature, and the composition discussed above, are inconsistent with iron or stony micrometeorites and other cosmic spherules, which are generally either chondritic or differentiated and do not show evidence of volatile degassing (24).

A key observation is the presence of amorphous, occasionally vesicular, nearly pure SiO2 glass (lechatelierite) inclusions within the microtektites (Fig. 3). Lechatelierite occurrence is confirmed by energy-dispersive x-ray spectroscopy (EDS) and micro-Raman spectroscopy in cross sections of spherules from Wilson Lake B, Millville, and Hole 1051B (see fig. S1B). EDS was used to identify inclusions that are >98% SiO2, and Raman spectra from several of these are consistent with that of fulgurite (lightning-induced lechatelierite). Lechatelierite forms from melting of quartz and quenching to quartz glass, which generally occurs at temperatures >1750°C (4, 5). This indicates that the P-E spherules formed at temperatures at least this high, which excludes any known terrestrial volcanism as their origin. Melting of crustal silicates by impact sufficient to produce lechatelierite occurs at shock pressures of ~30 to 50 GPa.

**Fig. 1.** Stratigraphic distribution of P-E spherules. (A to C) Stratigraphic distribution of P-E spherules from Millville (A), Wilson Lake B (B), and ODP Hole 1051B (C). Millville and Wilson Lake B core depths are in drilling units (du) of decimal feet; core depth in Hole 1051B is in meters below seafloor (mbsf). The carbon isotope composition of bulk carbonate from Millville and Wilson Lake B is from Wright and Schaller (J0), and Hole 1051B is from Katz et al. (J11). (D) Map showing location of the New Jersey (NJ) margin locations, including exposure in Medford, NJ, compared to that of Site 1051, Blake Nose. At both Wilson Lake B and Millville, the spherules are found in the basal Marlboro Clay Formation. Because the CIE has not been defined stratigraphically at the equivalent exposure of the Marlboro Clay at Medford, the stratigraphic placement of the spherules recovered there is only qualitative with respect to the P-E boundary (J12).
(25, 26). This observation implies that the target rock, although not necessarily silica-rich, at least had silt-sized or larger quartz grains that were melted on impact.

We have identified a quartz inclusion in a microtektite from Millville that has Raman spectral characteristics indicative of shocked quartz (Fig. 3D and fig. S21). In particular, the relaxation of the spectral peak corresponding to SiO$_2$ bond-bending vibration from 464 to 460 cm$^{-1}$ wave numbers observed in the Millville inclusion is consistent with observations from synthetic quartz experimentally shocked to peak pressures of 25.8 GPa (27) (fig. S21) and is considered diagnostic. Other characteristic spectral shifts unique to shocked quartz also match the Millville grain. An extensive search for individual shocked quartz among the abundant shelf quartz grains will be the subject of future investigation.

A subset of spherules shows quench-crystallization textures of clinopyroxene (Raman spectra most closely matching augite or diopside; see fig. S20), in an otherwise glassy matrix (Fig. 3), typical of microkrystites (3, 28). The feathered, dendritic, and chainlike textures of the P-E microkrystites are classic rapid-crystallization high-temperature quench features of impact glasses with lower silica content from other strewn fields (12). Examples are clinopyroxene (cpx) microkrystites of the upper Eocene spherule layer (17, 29, 30), attributed to the Popigai impact event (13, 15, 31), and the Cretaceous-Paleogene (K-Pg) spherules (also discussed below) (32). Among P-E microkrystites, vesicles are present but less common. Iron-rich inclusions are found in a few of the microkrystites and microtektites (fig. S16). These inclusions are generally noncrystalline and round or stringy. The presence of vesicles and lack of Ni-rich spinels in the P-E microkrystites is consistent with melt ejecta.

Within a given P-E spherule, the crystallite chemistry is close to the surrounding matrix glass. The distribution of major oxide chemistries is also similar to, but distinct from, the Late Eocene cpx spherules, which, like the P-E spherules, have high CaO contents (averaging only about 10 wt %, which is substantially lower than the P-E spherules). However, unlike the P-E spherules, the Late Eocene spherules have compositions much higher in SiO$_2$ and Na$_2$O and lower in Al$_2$O$_3$ and FeO (18, 30, 35). The P-E microtektites and microkrystites are roughly equally represented at each of the three sites (Wilson Lake B, Millville, and Hole 1051B).

Impact ejecta, including spherules, are associated with the 66-Ma K-Pg boundary on the New Jersey coastal plain, such as in the Bass River core (34). For the following reasons, we find it implausible that the K-Pg spherules were reworked and redeposited at the P-E boundary. First, the chemistry of the P-E spherules differs significantly from those found at K-Pg boundary sections, with the latter being much higher in silica content (40 to 68%) (25). Newly discovered deposits of unaltered microtektites and microkrystites at a K-Pg ejecta layer in Gorgonilla Island, Colombia (32), have much higher SiO$_2$ and slightly lower Al$_2$O$_3$ than the P-E spherules, and Al$_2$O$_3$ generally

![Fig. 2. Electron backscatter (15 kV) images of representative P-E spherules from Hole 1051B, Wilson Lake B, and Millville cores and the Medford exposure. Selected color micrographs are shown as insets. (A) Microtektite with a surface pit from ODP Hole 1051B (512.75 mbsf). (B) Microkrystite from ODP Hole 1051B (512.75 mbsf). Round features on surface of microkrystite are vesicles and small accretionary material. (C) Teardrop-shaped glass spherule from Millville 898.8 drilling units (du); inset is photomicrograph of same object. (D) Microkrystite with surface pit (spallation or microcrater?) from Millville (898.8 du). Dark round surface features are exposed vesicles. (E) Broken drop-form or dumbbell from Wilson Lake B (366.2 du). (F) Microtektite from Wilson Lake B (365.9 du), with a smaller spherule accreted to the side. This nature of deformation is typical of accretion occurring in partially molten state (compare to Fig. 3, E and F). (G) Microtektite with surface microcrater from the Medford, NJ Exposures. Inset shows several other forms found at Medford, which allows us to exclude drilling contamination as a source of the spherules. See table S4 for abundances. (H) Typical microkrystite from Wilson Lake B (366.2 du).]
increases with increasing silica content. These K-Pg spherules are also several times more enriched in Na₂O (average ~3%) compared with the P-E spherules (0.2%) and have about half as much FeO (5.5% in the K-Pg versus 10% in the P-E). The presence of conchoidal fracture patterns in the P-E spherule glass is also inconsistent with surficial exposure and chemical weathering (e.g., Fig. 2 and fig. S2, E and F). The K-Pg ejecta are not preserved as glassy spherules on the coastal plain and are usually diagnostically altered; furthermore, they were buried by 20 to 30 m of sediments by the time of the P-E boundary (36) and would have been unlikely to survive unaltered on the terrestrial surface for 10 million years before a redeposition event at the P-E boundary. Indeed, the occurrence of the P-E microtektites in a discrete time-correlative stratigraphic layer at three separate locations separated by >1000 km, two from the New Jersey shelf and one in the open ocean (Blake Nose) (Fig. 1), makes a required focused reworking of relatively rare K-Pg spherules highly improbable. In addition, if the spherules were reworked, they would be accompanied by other material of the same age, such as K-Pg microfossils, which would have been much more abundant than spherules. Also, the discovery of spherules at the Medford outcrop section (Fig. 2) that are identical in morphology and chemistry to those at Millville, Wilson Lake B, and Hole 1051B confirms that they are not an artifact of drilling disturbance.

We conclude that the spherule layer at Wilson Lake B, Millville, and ODP Site 1051 is best explained as air-fall ejecta (microtektites and microkrystites) generated by an impact close to the time of the P-E boundary. Ejecta fallout occurs over hours or days, implying that the spherule layer provides the most precise time-correlative feature for the P-E boundary among the three sites. The unprecedented high stratigraphic resolution of the Wilson Lake B and Millville sites, where the onset of the CIE is recorded over meters where the onset of the CIE is recorded over meters, compared with only a few centimeters in open ocean sections (e.g., ODP Sites 1051 and 690 (37)), provides the potential for temporal resolution that has not been heretofore possible and will surely be the target of important future work. Importantly, the virtually instantaneous time line provided by the microtektite horizon provides an opportunity to delineate the detailed sequencing of sedimentation and carbon cycling at a critical juncture in Earth's climate history.

For example, the apparent variable offsets between the microtektite horizon and inflections in the bulk sediment δ¹³C curves at Wilson Lake B and Millville, compared to the step function in foraminiferal δ¹³C (Fig. S22), must be viewed in this unfamiliar level of ultrahigh resolution and inspire greater scrutiny of the source(s) of the bulk carbon isotope excursion and its relation to the actual sequence of events at the P-E boundary.

It is worth noting that an Ir anomaly has been identified at a P-E boundary section at Zumaya, Spain (38), although it has been interpreted as volcanic in origin (39), and in flysch deposits from Slovenia (40); however, a large iridium anomaly is

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**Fig. 3.** Electron backscatter (15 kV) images of microtektites and microkrystites from polished cross sections. Scale in micrometers noted by white bars (L, lechatelierite inclusions; cpx, clinopyroxene microlites; V, vesicle). (A) Typical internal structure of a glassy microtektite from Wilson Lake B (365.5 du). (B) Teachdrop microkrystite from Millville (898.5 du). Note quartz and lechatelierite inclusion at edge. (C) Quench-form crystallite morphologies in a microkrystite from Millville (898.5 du). Note “chain form” and dendritic structures of cpx crystals, typical of rapid quenching from high temperatures. (D) Enlargement of grain inclusion in (B) showing lechatelierite around a quartz grain. Raman spectra from this quartz grain show characteristic vibrational shifts associated with shocked quartz (see fig. S21). Also note fine-grained dendritic cpx microlites at edges of droplet. (E) Vesicular lechatelierite accretion/inclusion surrounded by clinopyroxene microlites from Millville (898.8 du). Light-colored inclusions are iron-rich crystallites (see fig. S16 for details). (F) Microtektite showing accretion of a grain or an exposed inclusion with lechatelierite around it, from Millville (898.8 du). (G) Close-up of microkrystite showing dendritic and feather-form crystallites of clinopyroxene from Hole 1051B (512.75 mbsf). (H) Typical microtektite with small lechatelierite inclusion from Hole 1051B (512.75 mbsf).
not necessarily associated with all major impacts [e.g., the Chesapeake impact (47)]. These findings will motivate a search for impact ejecta at other sites to define the geographic footprint of the P-E strewn field, which will ultimately constrain the currently unknown location of an impact crater.

REFERENCES AND NOTES

12. All primary data are available as supplementary materials on Science Online.

PROTEIN EVOLUTION

Evolution of protein phosphorylation across 18 fungal species

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Living organisms have evolved protein phosphorylation, a rapid and versatile mechanism that drives signaling and regulates protein function. We report the phosphoproteomes of 18 fungal species and a phylogenetic-based approach to study phosphosite evolution. We observe rapid divergence, with only a small fraction of phosphosites conserved over hundreds of millions of years. Relative to recently acquired phosphosites, ancient sites are enriched at protein interfaces and are more likely to be functionally important, as we show for sites on H2A1 and eIF4E. We also observe a change in phosphosite motif frequencies and kinase activities that coincides with the whole-genome duplication event. Our results provide an evolutionary history for phosphosites and suggest that rapid evolution of phosphorylation can contribute strongly to phenotypic diversity.

Divergence in transcriptional regulation and gene expression is often considered the main driver of phenotypic variation across species (1). Comparative gene expression and chromatin immunoprecipitation sequencing studies show that transcriptional regulation has diverged very quickly (2). Similar analysis of protein posttranslational regulation has lagged due to methodological limitations. Now, mass spectrometry (MS) allows comprehensive
Supplementary Materials for

**Impact ejecta at the Paleocene-Eocene boundary**

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This PDF file includes:

- Materials and Methods
- Supplementary Text
- Figs. S1 to S22
- Tables S1 to S6
- References
Materials and Methods

Samples of ~30-40 g dry weight (where available) were taken below and through the onset of the CIE from the Wilson Lake B and Millville cores, avoiding any portions showing evidence of drilling mud injection. At ODP Hole 1051B, we used samples taken for foraminiferal assemblages and isotopes (/I), with sample sizes between 2 and 20 g. Sediment samples were soaked in DI, washed through a 63μm sieve and dried in an oven at 50°C. Spherules were picked manually from the >63 μm size fraction and reported as number of spherules per gram of dry (unwashed) sediment. The shelf sediments from the Marlboro clay are dominated by kayolinite and illite clays with silts in the fine fraction, and glauconite, quartz and mica in the >63 μm size fraction (42). At site 1051 the fine fraction is mostly coccoliths, with radiolaria and foraminifera and occasional clastic grains (quartz) in the course fraction. To minimize potential observational bias, samples were randomized by a disinterested third party, and those picking were not aware of which samples were likely to encompass the spherule layer and searches of each sample (including those where no glassy spherules were recovered) were allotted up to 50 hours. Spherules were mounted on glass slides for micro-laser Raman spectroscopy using the Bruker 532 nm green laser system at RPI. Separate grain mounts of glass particles were carbon-coated for analysis by SEM or ion microprobe. Selected samples were mounted in epoxy and polished using a 0.03 μm grit to reveal their cross sections, which were also analyzed by Raman spectroscopy. Energy-dispersive X-ray spectroscopy (EDS) measurements were made using a Bruker x-flash 5010 detector (with Be window) mounted on the Cameca SX-100 ion microprobe at RPI for both grain and epoxy mounts. Quantitative major element chemistries were determined by companion EDS measurements of standard glasses and minerals. At least two EDS spectra were generated for each specimen and averaged after removal of background using Bruker’s Esprit software. Chemistries were normalized to major oxides assuming stoichiometry.

Supplementary Text

The Paleocene-Eocene boundary at Wilson Lake and implications for other coastal plain sections

The Paleocene-Eocene boundary is defined by a global decrease in δ13C going up section (7) recorded in marine settings by foraminifers, bulk sediments (dominated by nanofossils) and organic matter (37, 43-44). Terrestrial sections record large δ13C decreases in paleosols, mammalian teeth, and terrestrial and lacustrine organic matter (45, 46). At most sites, the onset occurs over thin stratigraphic intervals with few if any transitional δ13C values, allowing unambiguous placement of the P-E boundary. However, the Atlantic coastal plain from Virginia to New Jersey yields expanded records of the onset with many intermediate values, indicating a dramatic increase in sedimentation rates and an unprecedented chronicle of events across the P-E boundary (10).

At Wilson Lake, the CIE onset differs in placement depending whether the inorganic δ13C record from bulk sediments is used versus other δ13C recorders, with the bulk decrease starting ~1.2 m below the organic δ13C measurements (47). The scarcity of foraminifers in the coarse-grained, glauconitic, sandy silts (Vincentown Formation in
New Jersey and Aquia Formation in Maryland/Virginia) below the Marlboro Clay make the bulk records attractive for correlation and boundary placement. Figure S13 shows the relationships between the bulk and foraminiferal $\delta^{13}C$ at Wilson Lake A (48) along with the Sluijs et al. (47) pick for the onset of the CIE (and hence the P-E boundary) based on dinoflagellate $\delta^{13}C$ values. We agree with Sluijs et al. (47) on this placement, giving more weight to the foraminiferal and dinoflagellate $δ^{13}C$ records over the bulk carbonate record. This also corresponds to the change in lithology from glauconitic sandy silts to the clayey silts of the Marlboro Clay (49) and the major decrease in %CaCO$_3$ that is common at the onset of the CIE (10, 47).

The bulk $δ^{13}C$ records show that the nearby (within ~50 meters) Wilson Lake –A and –B core depths can be precisely correlated, but are offset by ~1.5 m (~5 feet) with common features being deeper in the B-Hole. Mapping the Sluijs et al. pick for the P-E boundary based on dinoflagellate $δ^{13}C$ values from the A-Hole to the B-Hole places it at ~111.55 m (366 ft) in the B-Hole, which matches the peak concentrations in spherules (Figure S22). The process(es) that may have produced the lead in bulk $δ^{13}C$ relative to the other $δ^{13}C$ records across the onset of the CIE will be the subject of further study.
Fig. S1.
Light Micrographs of spherules from Wilson Lake B and Millville. A. Microtektites and microkrystites from Wilson Lake B (366.2 du), B. Enlargement of various microtektites from Wilson Lake B (366.2 du, drilling units, in decimal feet), C. Microtektites and microkrystites from Millville (898.8 du), D. Enlargement of spherules and broken rotational form from Millville (898.8 du).
**Fig. S2**

Electron backscatter (15 kV) image of a microtektite from Wilson Lake B (365.9 drilling units (du)). Images are taken using a CAMECA SX100 on the ion microprobe at RPI. Scale in microns noted by white bars, standard contrast scale with saturation is shown to the right of each image.
Fig. S3
Microtektite from Wilson Lake B (365.9 du).
Fig. S4
Microtektite from Hole 1051B (512.75 mbsf).
Fig. S5
Rotational form microtektite from Medford outcrop (3.81 cm).
Fig. S6
Rotational form microtektite from Millville (898.8 du).
Fig. S7
Microkrystite from Medford outcrop with a microcrater.
Fig. S8
Electron backscatter (15 kV) image of a polished cross-section of a microtektite from Millville (898.8 du). Inclusion on top is probably lechatelierite. Images are taken using a CAMECA SX100 on the ion microprobe at RPI.
Fig. S9
Microtektite from Wilson Lake B (265.9 du). Bright spots are dust particles.
Fig. S10
Microtektite from Hole 1051B (512.75 mbsf). Inclusion at bottom is probably lechatelierite, dark spot at center is a vesicle.
Fig. S11
Close-up of Microkrystite from Wilson Lake B (365.5 du) showing lath-like and feather-form crystallites of clinopyroxene (cpx).
Fig. S12
Blown-up image of a microkrystite from Millville (898.5 du) showing dendritic clinopyroxene crystallites and lechatelierite inclusions (both at right) side.
Fig. S13
Microkrystite from Wilson Lake (365.5 du) showing feathered texture of cpx crystals.
Fig. S14
Energy-dispersive X-ray spectroscopy (EDS) spectrum of a representative polished cross-section of a microtektite matrix from Hole 1051B (512.75 mbsf). EDS measurements are made using the Bruker x flash 5010 detector (with Be window) and background corrected on the Esprit v.1.8 software. Y-axis in cps/eV.
Fig. S15
EDS spectrum of a representative lechatelierite inclusion in a microtektite from Millville (898.5 du).
Fig. S16
Backscatter image (bottom) and EDS spectrum (top) of an iron-rich inclusion in a microtektite from Millville (898.8 du). The EDS measurement is from the inclusion on the right. Note the large vesicle near the center of the spherule.
Fig. S17
Micro-laser Raman spectra of representative microtektite matrices. Spectra are collected using a Bruker 532 nm green laser system at RPI and are background corrected.
Fig. S18
Background corrected Raman spectra of representative lechatelierite inclusions found in spherules compared to the spherule matrix, lechatelierite inclusions in fulgurite, SiO$_2$ glass, and quartz. Similar peaks in the 1250-1450 wavelength region in the lechatelierite spectrum are effects of the spherule matrix, see Fig. S19 for location of analyses.
Fig. S19
Electron backscatter image of a polished spherule (Millville 898.8 du) example of one used to identify the phases by Raman spectroscopy, above. Lechatelierite inclusion (L), matrix (M), clinopyroxene crystalites (cpx), crystalline α-quartz (α-quartz), and vesicles (V) are labeled.
Fig. S20
Raman spectra of representative clinopyroxene crystallites as compared to augite, diopside, and ferrosilite standards. Spectra are background corrected.
Fig. S21
A. Background corrected Raman spectrum collected from a polished microkrystite containing crystalline α-quartz (Millville 898.5 du) and an unshocked quartz standard. The black dashed lines note shifts in various vibrational modes between an unshocked quartz grain and the quartz inclusion in the microkrystite. These measurements are compared to the spectra for quartz shocked to peak pressures of 25.8 GPa (26), which are remarkably consistent with those from the Millville inclusion. The inset shows the characteristic 464 to 450 wavenumber vibrational relaxation associated with shocked quartz. Data from McMillan et al (26) were digitized from a paper copy. We also show a RRUFF database spectrum for Coesite for reference. B. is an electron backscatter image of the polished microkrystite containing the crystalline shocked quartz inclusion (Millville 898.5 du) measured, M = matrix, cpx = clinopyroxene microlites, L = lechatelierite.
Fig. S22
Correlation of $\delta^{13}$C records from Wilson Lake A and B Holes during the onset of the CIE. The vertical black line at 360.9 ft in Wilson Lake A shows the location of the CIE onset determined by Sluijs et al. (50) using a dinoflagellate $\delta^{13}$C record. Also shown are two planktonic foraminiferal values from Wilson Lake A (51), which are similar to regional $\delta^{13}$C values for each genus in pre-CIE intervals of other sites (red and green shaded boxes) (51). At the bottom of the graph, abundances of spherules found in the B-Hole are shown, and closely match the Sluijs et al. (50) pick for the P-E boundary based on the onset of the CIE.
<table>
<thead>
<tr>
<th>mbsf</th>
<th>Section</th>
<th>Sample Interval (cm)</th>
<th>Pre-Washed Weight (g)</th>
<th>Post-Washed Weight &gt;63μm (g)</th>
<th>Post-Washed Weight 45-63μm (g)</th>
<th>Spherules &gt;63μm</th>
<th>#/gram</th>
</tr>
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<tbody>
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<td>0.01</td>
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<td>0.69</td>
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<td>0.02</td>
<td>0</td>
<td>0</td>
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<td>1.51</td>
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**Table S1.**

Hole 1051B spherule concentrations (#/gram) and corresponding sample depth.
<table>
<thead>
<tr>
<th>Sample (ft)</th>
<th>Pre-Washed Weight (g)</th>
<th>Post-Washed Weight &gt;63μm (g)</th>
<th>Post-Washed Weight 45-63μm (g)</th>
<th>Spherules &gt;63μm</th>
<th>#/gram</th>
</tr>
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<tbody>
<tr>
<td>364.35</td>
<td>4.46</td>
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<td>0.07</td>
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<td>5.53</td>
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<td>0.09</td>
<td>0</td>
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<td>365.10</td>
<td>6.26</td>
<td>0.63</td>
<td>0.11</td>
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<td>4.86</td>
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<td>365.50</td>
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<td>0.21</td>
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<td>1.35</td>
<td>0.14</td>
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<tr>
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<td>1.23</td>
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<td>3.81</td>
<td>1.67</td>
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<td>373.30</td>
<td>5.52</td>
<td>2.16</td>
<td>0.36</td>
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</table>

**Table S2.**

Wilson Lake B spherule concentrations (#/gram) and corresponding sample depth.
Table S3.
Millville spherule concentrations (#/gram) and corresponding sample depth.

<table>
<thead>
<tr>
<th>Sample (ft)</th>
<th>Adj. Sample (ft)</th>
<th>Pre-Washed Weight (g)</th>
<th>Post-Washed Weight &gt;63μm (g)</th>
<th>Post-Washed Weight 45-63μm (g)</th>
<th>Spherules &gt;63μm</th>
<th>#/gram</th>
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<tbody>
<tr>
<td>898.20</td>
<td>897.70</td>
<td>8.36</td>
<td>0.76</td>
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<td>898.50</td>
<td>897.98</td>
<td>8</td>
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<td>0.25</td>
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<tr>
<td>898.80</td>
<td>898.26</td>
<td>8.21</td>
<td>1.17</td>
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<td>2.07</td>
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<td>899.40</td>
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</table>
**Table S4.**

Medford Exposure spherule concentrations (#/gram) and corresponding sample depth (inches) from a short (50 cm) push-core taken at the exposure. Sample depths measured from the bottom of the core.

<table>
<thead>
<tr>
<th>Sample Depth (in from Bottom)</th>
<th>Sample Depth (cm from Bottom)</th>
<th>Pre-Washed Weight (g)</th>
<th>Post-Washed Weight &gt;63μm (g)</th>
<th>Spherules &gt;63μm</th>
<th>#/gram</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.5</td>
<td>0-1.5</td>
<td>3.81</td>
<td>62.1</td>
<td>1.22</td>
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<tr>
<td>3</td>
<td>2.0-3.0</td>
<td>7.62</td>
<td>61.1</td>
<td>1.09</td>
<td>0</td>
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<tr>
<td>4.5</td>
<td>3.5-4.5</td>
<td>11.43</td>
<td>63.75</td>
<td>1.28</td>
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<td>16.5</td>
<td>15.5-16.5</td>
<td>41.91</td>
<td>64.67</td>
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</table>
Table S5.
Weight % H₂O content determined by FTIR from polished sections of spherules.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Integrated Molar Absorption Coefficient</th>
<th>Weight % H₂O**</th>
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</thead>
<tbody>
<tr>
<td>Medford 3.81 cm (Microtektite)</td>
<td>32000 ± 3200</td>
<td>0.0228 ± 0.00173</td>
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<tr>
<td>Wilson Lake B 365.9 du A (Microtektite)</td>
<td>32000 ± 3200</td>
<td>0.0234 ± 0.000782</td>
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<td>Wilson Lake B 365.9 du B (Microkrystite)</td>
<td>32000 ± 3200</td>
<td>0.00988 ± 0.00253</td>
</tr>
</tbody>
</table>

* Silver and Stolper (1985)
** 1 o standard deviation of several spectra (3-7) collected on a single microtektite/microkrystite
Table S6.

Major oxide chemistries determined by EDS of the matrix from either grain mount or polished spherules from sites 1051 (512.75 mbsf), Millville (MV), Wilson Lake B (WL), and Medford (MD). Spherules are separated by microtektites and microkrystites and sorted by SiO$_2$ content.
References


12. All primary data are available as supplementary materials on *Science* Online.


