**Fig. 3.** Comparative plot of active houses of giant larvaceans (blue line) and discarded sinkers (red line) versus depth, in square meters of area swept. The data are derived from a 10-year time series of quantitative video transects at depth intervals between 100 and 1000 m ($n = 679$ transects). With an average sinking rate of 800 m day$^{-1}$, the difference between the integrated areas beneath the curves indicates that these animals produce a new house each day (24).

Rapid Acidification of the Ocean During the Paleocene-Eocene Thermal Maximum


The Paleocene-Eocene thermal maximum (PETM) has been attributed to the rapid release of $\sim 2000 \times 10^9$ metric tons of carbon in the form of methane. In theory, oxidation and ocean absorption of this carbon should have lowered deep-sea pH, thereby triggering a rapid ($\sim 10,000$-year) shoaling of the calcite compensation depth (CCD), followed by gradual recovery. Here we present geochemo-data from five new South Atlantic deep-sea sections that constrain the timing and extent of massive sea-floor carbonate dissolution coincident with the PETM. The sections, from between 2.7 and 4.8 kilometers water depth, are marked by a prominent clay layer, the character of which indicates that the CCD shoaled rapidly ($\sim 10,000$ years) by more than 2 kilometers and recovered gradually ($\sim 100,000$ years). These findings indicate that a large mass of carbon ($\sim 2000 \times 10^9$ metric tons of carbon) dissolved in the ocean at the Paleocene-Eocene boundary and that permanent sequestration of this carbon occurred through silicate weathering feedback.

During the Paleocene-Eocene thermal maximum (PETM), sea surface temperature (SST) rose by 5°C in the tropics and as much as 9°C at high latitudes (1–3), whereas bottom-water temperatures increased by 4° to 5°C (4). The initial SST rise was rapid, on the order of $\sim 10^4$ years, although the full extent of warming was not reached until some $\sim 30,000$ years (30 ky).
later (5). The most compelling evidence for greenhouse forcing is a coeval global carbon isotope excursion (CIE) of roughly −3.0 per mil (%e) in deep-sea cores (4). The pattern of the CIE—an initial rapid decrease (~20 ky) followed by a more gradual recovery (130 to 190 ky) (1, 6–8)—indicates the input of a large mass of isotopically depleted carbon into the ocean and atmosphere. Quantitatively, methane hydrates, with a mean δ¹³C of ~−67%, appear to be the most plausible source of this carbon (9). For example, only ~1200 × 10⁹ metric tons of carbon (GtC) of biogenic methane would be required to produce a CIE of 2.5% (10, 11). Thermogenic methane has been implicated as well (12), although the mass required to produce the CIE would be roughly double that of the biogenic methane.

Regardless of its source, the released methane was rapidly oxidized to CO₂. Subsequent oceanic dissolution of this CO₂ would alter ocean carbon chemistry, principally by lowering the pH and carbonate ion content of seawater. These changes would be partially neutralized by a transient rise in the level of the lysocline and calcite compensation depth (CCD) (13), resulting in the widespread dissolution of sea-floor carbonate. Eventually, the CO₂ would be sequestered and ocean carbonate chemistry would be restored, primarily through chemical weathering of silicate rocks (10). The extent and duration of lysocline/CCD shoaling and subsequent recovery would depend largely on the source, mass, and rate of carbon input. For example, modeling of a 1200-GtC input over 10 ky produces a lysocline shoaling of 300 m (less in the Pacific) with a recovery time of ~40 ky (10). Such changes in [CO₂] should produce distinct patterns in pelagic carbonate sedimentation and lithology, characterized by an abrupt transition from carbonate-rich sediment to clay, followed by a gradual recovery to carbonate. Moreover, the clay layer should increase in thickness with increasing water depth.

Clay or low-carbonate layers coincident with the PETM were previously identified in several deep-sea cores and land-based marine sections (14–16). However, these sections, which are either geographically isolated or not completely recovered, or both, are inadequate for constraining CCD variations and for testing the methane hypothesis. Ocean Drilling Program (ODP) Leg 208 was designed to recover an array of pelagic cores spanning the Paleocene-Eocene (P-E) boundary over a broad depth range. The primary drilling target was the Walvis Ridge, in the southeastern Atlantic (fig. S1), where the Deep Sea Drilling Project (DSDP) Leg 74 rotary cored portions of the P-E boundary sequence near the base and summit of the ridge (sites 527 and 525) (17). By using advanced piston coring in multiple offset holes at five sites (1262, 1263, 1265, 1266, and 1267), Leg 208 successfully recovered stratigraphically complete and undisturbed upper Paleocene-to-lower Eocene successions at four of five sites between 2.7 and 4.8 km water depth (18). At each site, the P-E boundary sequence was characterized by an abrupt transition from carbonate-rich ooze to a dark
Fig. 2. Bulk sediment carbon isotope records for holes 1262A, 1263C/D, 1265A, 1266C, and 1267B plotted versus MCD. Also plotted are nanofossil horizons (N1 to N4, arrows in red) for holes 1262B and 1263C/D (20). Data for ODP site 690 (22) are plotted to the far left versus meters below the sea floor (MBSF). Lines of correlation are based on inflections in the carbon isotope (A to G above the P-E boundary, –A below), Fe/Ca, and magnetic susceptibility (MS) records (20). vPDB, Vienna PeeDee Belemnite.

red “clay layer,” which then graded back into ooze (Fig. 1 and table S1). Carbonate content was <1% weight percent (wt %) in the clay layers, and >80 and 90 wt % in the underlying and overlying oozes, respectively; the only exception was site 1265, where the basal portion of the clay layer was not recovered. The thickness of the clay layers increased with depth, from 5 cm at the shallowest site (1263) [2717 m; paleodepth ~1500 m (19)] to 35 cm in the deepest site (1262) [4755 m; paleodepth ~3600 m (19)] (Fig. 1). The benthic foraminiferal extinction horizon, which is characterized by the disappearance of long-lived Paleocene species and a rapid drop in diversity, occurred at the base of the clay layer in each site (19).

Bulk sediment carbon isotope ($\delta^{13}C$) records were constructed at 1- to 5-cm resolution for each boundary section (table S2) (20). Each record is marked by a decrease in $\delta^{13}C$ at the base of the clay layer, followed by gradual recovery. Minimum carbon isotope values within the clay layer are not uniform, but increase from the shallowest to the deepest site (minimum of –0.9 and 0.0% at sites 1263 and 1262, respectively), a feature we attribute to truncation by dissolution and the presence of residual pre-exursion calcite (21). Also, the base of the CIE differs across sites, occurring in two steps at site 1263 and in a single step at the deeper sites. As a result, the excursion layer, from the onset of the CIE to the point of full recovery (i.e., stability), decreases in thickness from 2.1 m at site 1263 to 1.0 m at 1262.

In this spatially tight array of sites, the production and export of carbonate and the accumulation of clay should be similar at any given time, leaving dissolution as the major process that drives differences in carbonate accumulation between sites. We can therefore infer from the weight % carbonate and carbon isotope data that rapid shoaling of the lysocline/CCD occurred, followed first by a more gradual descent or recovery of the CCD and then by the recovery of the lysocline. The duration of the lysocline/CCD descent from the shallowest to the deepest sites was estimated by first correlating several key inflection points in the carbon isotope records (Fig. 2, tie points A to G), as well as in the Fe concentration and bulk magnetic susceptibility (MS) records (Fig. S2). The tie points, particularly E and F, were then verified with biostratigraphic data (table S3) (20). We then correlated the site 1263 carbon isotope record to that of south Atlantic ODP site 690 (22), which has an orbitally derived age model (8), and ordinated the weight % carbonate and isotope data for each site within that age model (Fig. 3 and table S4). An alternate age model based on 3He exists for site 690 (23), but the two models are roughly similar for the initial 100 ky of the PETM; thus, the choice of model makes little difference in our interpretation of events up to that point. The greatest uncertainty in the site-to-site correlations and age estimates is in the basal portion of the clay layer, where the carbon isotope and other records are compromised by dissolution. The correlations (Fig. 1, tie points D to G) are most reliable in the recovery interval where the weight % carbonate is higher and the ocean $\delta^{13}C$ is rapidly shifting.

Given these age constraints, the CCD is inferred to have shoaled more than 2 km within a few thousand years (Fig. 3). Recovery was gradual, with the CCD descending to the shallowest site (1263) within ~10 to 15 ky of the CIE onset and to the deepest site (1262) within ~60 ky. By +110 ky, carbonate content had fully recovered. This pattern of change, particularly the recovery, has important implications. According to theory, the initial uptake of CO₂ and buffering should occur mainly via deep-sea calcite dissolution, but eventually, chemical weathering of silicate rocks takes over accelerating the flux of dissolved ions (including HCO₃⁻) to the ocean, thereby increasing [CO₂] and the rate of calcite accumulation (24). The distribution of carbonate between +60 and +100 ky indicates that the CCD had descended, but the lysocline was still shallow and the deep sea was largely undersaturated. The percentage of CaCO₃ continued to increase, and by +110 kyr, it had reached 90% over the entire
The mass of CO₂ required to shoal the CCD to <1 km water depth would be substantial. In a series of simulations with an ocean/sediment carbon-cycle model designed to evaluate the ocean-buffering capacity in response to a range of anthropogenic CO₂ fluxes, 4500 GtC was required to terminate carbonate accumulation over the entire ocean (26).

For the PETM, the release of >4500 GtC would be more consistent with the magnitude of global temperature rise (2, 3, 9). Such a large mass of carbon, however, would require a reevaluation of the source of carbon and its isotopic composition. With bacterially produced methane at ~60%, the total input from hydrates is limited by the δ¹³C excursion to ~2000 GtC (10). To increase the mass of carbon added while adhering to the isotope constraints requires the input of isotopically heavier carbon, such as thermogenic CH₄/CO₂ (~30 to ~20%) or oxidation of organic carbon (standing or stored) (~20%) (34). In this regard, recent documentation of an unusual concentration of upper Paleocene fluid/gas deep conduits associated with volcanic intrusions in the North Atlantic (12) merits additional attention. An alternative explanation, that the magnitude of the marine CIE has been greatly underestimated because of dissolution or damping by pH effects, seems unlikely given the constraints provided by continental isotope records (35). Finally, proximity to where carbon (CO₂ or CH₄) enters the deep sea via circulation will dictate where neutralization by carbonate dissolution is most intense (36). For example, severe dissolution in the Atlantic may indicate direct input of methane into bottom waters entering this basin.

Excessive carbonate undersaturation of the deep ocean would likely impede calcification by marine organisms and therefore is a potential contributing factor to the mass extinction of benthic foraminifera at the P-E boundary. Although most plankton species survived, carbonate ion changes in the surface ocean might have contributed to the brief appearance of weakly calcified planktonic foraminifera (6) and the dominance of heavily calcified forms of calcareous algae (37). What, if any, implications might this have for the future? If combustion of the entire fossil fuel reservoir (~4500 GtC) is assumed, the impacts on deep-sea pH and biota will likely be similar to those in the PETM. However, because the anthropogenic carbon input will occur within just 300 years, which is less than the mixing time of the ocean (38), the impacts on surface ocean pH and biota will probably be more severe.

References and Notes
13. The lysocline, also referred to as the calcite saturation horizon, represents the depth in the ocean where the carbonate ion concentration falls below the saturation level (currently ~4 km in the South Atlantic). Carbonate accumulation can occur below this level if the flux of carbonate to the sea floor exceeds the rate at which it dissolves. The depth at which dissolution is greater than the flux and where carbonate does not accumulate defines the CCD.

Fig. 3. Bulk sediment δ¹³C and weight % carbonate content (wco2/WTOTAL × 100) plotted versus age for ODP sites 1262, 1263, 1265, 1266, and 1267. Age (ky) relative to the P-E boundary is plotted on the left axis and absolute age (Ma) along the right. Age models (table S4) are based on correlation to site 690 (8) using the carbon isotope stratigraphy as verified with the nannofossil events in Fig. 2 and with the Fe and M5 cycles in fig. S2. Transferring the 1263 age model to deeper sites with carbon isotopes could only be achieved where sufficient carbonate was present. Ages within the clay layers for sites 1266, 1267, and 1262 were derived through linear interpolation from tie points E and A. Paleodepths (~55 Ma) are provided for sites 1263 (1500 m), 1266 (2600 m), and 1262 (3600 m). Key events in the evolution of South Atlantic carbonate chemistry were: (i) the rapid drop in content to ~1% for all sites with the exception of site 1265, where the lowermost Eocene is absent (marked I); (ii) the return of the CCD to site 1263 roughly 5 ky after the excursion (marked II); (iii) the return of the CCD to site 1262 at 60 ky (marked III); and (iv) the lysocline descending to a point below the deepest site at 110 ky after the excursion (marked IV). PEB, Paleocene-Eocene boundary.
20. Materials and methods are available as supporting material on Science Online.

21. The initial phase of dissolution would involve Paleocene sediments already present on the sea floor. As such, the base of the clay layer, perhaps as much as a few centimeters in the deepest site, was deposited before the carbon isotope excursion. The traces of carbonates remaining must be a mixture of pre-excursion and excursion fragments that survived dissolution.


24. The weathering of silicates on land is generally represented by the following equation: CaSiO₄ + 2CO₂ + H₂O → 2HCO₃⁻ + Ca²⁺ + SiO₂. Ensuing precipitation of calcite from the bicarbonate (and carbonate) ions supplied by the above reaction is represented by this equation: HCO₃⁻ + Ca²⁺ → CaCO₃ + CO₂ + H₂O, so that there is a net uptake of one unit of CO₂ for each unit of silicate weathered.


The primary buffering capacity of the ocean is provided by the deep ocean and sea-floor sediments. Because the mixing time of the ocean is ~500 years, most of the anthropogenic CO₂ will accumulate in the atmosphere and surface ocean before it can be conveyed to the deep sea to be neutralized (27).

38. We thank the ODP Leg 208 Science Crew for their contributions and C. John and S. Bohaty for technical assistance. The ODP supplied samples. Supported by NSF grant EAR-0120727 to J.C.Z. and E.T. and by DFG grant no. Ro 1113/3 to U.R.

Supporting Online Material
www.sciencemag.org/cgi/content/full/308/572/1611/DC1
Materials and Methods
Figs. S1 and S2
Tables S1 to S4
References
21 December 2004; accepted 18 April 2005
10.1126/science.1109004

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**Photoinduced Plasticity in Cross-Linked Polymers**

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Chemically cross-linked polymers are inherently limited by stresses that are introduced by post-gelation volume changes during polymerization. It is also difficult to change a cross-linked polymer’s shape without a corresponding loss of material properties or substantial stress development. We demonstrate a cross-linked polymer that, upon exposure to light, exhibits stress and/or strain relaxation without any concomitant change in material properties. This result is achieved by introducing radicals via photocleavage of residual photoinitiator in the polymer matrix, which then diffuse via addition-fragmentation chain transfer of midchain functional groups. These processes lead to photoinduced plasticity, actuation, and equilibrium shape changes without residual stress. Such polymeric materials are critical to the development of microdevices, biomaterials, and polymeric coatings.

Cross-linked, gelled polymers have an “infinite” molecular weight and are described as thermosts, implying a network that cannot be melted or molded (1). This description is true for most chemically cross-linked polymers; however, several cross-linked networks are known to undergo bond cleavage or depolymerization at high temperatures or under various chemical or other treatments (2). Although such treatments are useful for recycling purposes, there is an associated degradation in the mechanical properties of the polymers. “Crack-healing” networks, such as those that use groups in the polymer backbone able to undergo thermoreversible Diels-Alder reactions (3), are able to relieve stress without mechanical degradation. However, this reaction must be performed at elevated temperatures, making it unsuitable in thermally sensitive applications such as dental composites. Internal stress buildup during polymerization is typical when shrinkage occurs. This stress decreases the ultimate mechanical properties of the cured polymer, which is highly detrimental in fields such as polymeric coatings, fiber-reinforced composites, and dental materials, or it may introduce birefringence, unwanted in optical materials. Additionally, given that the equilibrium shape of conventional cross-linked polymers is defined by the shape at gelation, stress relief would enable a material to be “molded” and subsequently destressed, allowing for arbitrary equilibrium shapes to be attained after cure.

We describe a covalently cross-linked network that is able to undergo photomediated, reversible cleavage of its backbone to allow chain rearrangement for rapid stress relief at ambient conditions without mechanical property degradation. The key to this reversible backbone cleavage is addition-fragmentation chain transfer. Reaction diffusion of radicals through the cross-linked matrix occurs initially by reaction of a radical with an in-chain functionality, forming an intermediate, which in turn fragments, reforming the initial functionality and radical. Allyl sulfides have been used as efficient addition-fragmentation chain transfer agents (4–6). This addition-fragmentation process alters the topology of the network, but the polymer chemistry and network connectivity remain unchanged. In the absence of radical termination events or other side reactions, the number of allyl sulfide groups, and hence network strands, remains unchanged (Scheme 1), although relaxation of the stresses in each bond is facilitated by the alternating cleavage and reformation reactions.

The monomers used to produce the networks are shown in Scheme 2. The base network studied was formed from a stoichiometric mixture of pentaerythritol tetra(3-mercaptopropionate) (PETMP) and triethyleneglycol divinyl ether (TEGDVE), which produces a rubbery network with a glass transition temperature (Tₘ) of about -25°C. This monomer system was modified by the addition of varying concentrations of the ring-opening monomer 2-methyl-1,7-methylene-1,5-dithiaocyclooctane (MDTO) (7) as a comonomer. Addition of a stoi-