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### Evidence for Upwelling of Corrosive "Acidified" Water onto the Continental Shelf

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The absorption of atmospheric carbon dioxide  $(CO_2)$  into the ocean lowers the pH of the waters. This so-called ocean acidification could have important consequences for marine ecosystems. To better understand the extent of this ocean acidification in coastal waters, we conducted hydrographic surveys along the continental shelf of western North America from central Canada to northern Mexico. We observed seawater that is undersaturated with respect to aragonite upwelling onto large portions of the continental shelf, reaching depths of ~40 to 120 meters along most transect lines and all the way to the surface on one transect off northern California. Although seasonal upwelling of the undersaturated waters onto the shelf is a natural phenomenon in this region, the ocean uptake of anthropogenic  $CO_2$  has increased the areal extent of the affected area.

ver the past 250 years, the release of carbon dioxide (CO<sub>2</sub>) from industrial and agricultural activities has resulted in atmospheric CO<sub>2</sub> concentrations that have increased by about 100 parts per million (ppm). The atmospheric concentration of CO<sub>2</sub> is now higher than it has been for at least the past 650,000 years, and is expected to continue to rise at an increasing rate, leading to pronounced changes in our climate by the end of this century (1). Since the beginning of the industrial era, the oceans have absorbed  $\sim$ 127  $\pm$  18 billion metric tons of carbon as CO<sub>2</sub> from the atmosphere, or about one-third of the anthropogenic carbon emissions released (2). This process of absorption of anthropogenic CO<sub>2</sub> has benefited humankind by substantially reducing the greenhouse gas concentrations in the atmosphere and minimizing some of the impacts of global warming. However, the ocean's daily uptake of 22 million metric tons of CO<sub>2</sub> has a sizable impact on its chemistry and biology. Recent hydrographic surveys and modeling studies have confirmed that the uptake of anthropogenic CO<sub>2</sub> by the oceans has resulted in a lowering of seawater pH by about 0.1 since the beginning of the industrial revolution (3–7). In the coming decades, this phenomenon, called "ocean acidification," could affect some of the most fundamental biological and geochemical processes of the sea and seriously alter the fundamental structure of pelagic and benthic ecosystems (8).

Estimates of future atmospheric and oceanic CO<sub>2</sub> concentrations, based on the Intergovernmental Panel on Climate Change (IPCC) CO<sub>2</sub> emission scenarios and general circulation models, indicate

that atmospheric  $CO_2$  concentrations could exceed 500 ppm by the middle of this century, and 800 ppm near the end of the century. This increase would

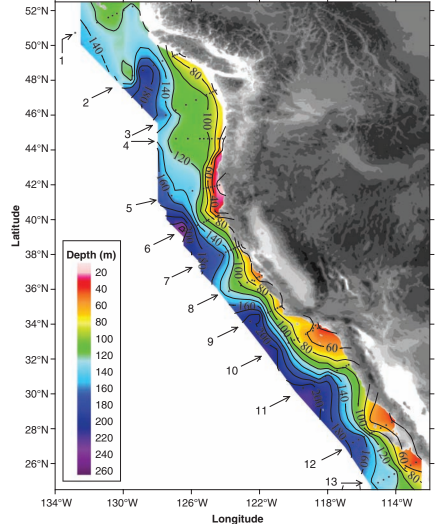
result in a decrease in surface-water pH of  $\sim 0.4$  by the end of the century, and a corresponding 50% decrease in carbonate ion concentration (5, 9). Such rapid changes are likely to negatively affect marine ecosystems, seriously jeopardizing the multifaceted economies that currently depend on them (10).

The reaction of  $CO_2$  with seawater reduces the availability of carbonate ions that are necessary for calcium carbonate (CaCO<sub>3</sub>) skeleton and shell formation for marine organisms such as corals, marine plankton, and shellfish. The extent to which the organisms are affected depends largely on the  $CaCO_3$  saturation state ( $\Omega$ ), which is the product of the concentrations of  $Ca^{2+}$  and  $CO_3^{2-}$  divided by the apparent stoichiometric solubility product for either aragonite or calcite:

$$\Omega_{\text{arag}} = [\text{Ca}^{2+}][\text{CO}_3^{2-}]/K'_{\text{sp}_{\text{arag}}}$$
(1)

$$\Omega_{\rm cal} = [{\rm Ca}^{2+}][{\rm CO_3}^{2-}]/K'_{\rm sp_{\rm cal}}$$
 (2)

where the calcium concentration is estimated from the salinity, and the carbonate ion con-



**Fig. 1.** Distribution of the depths of the undersaturated water (aragonite saturation < 1.0; pH < 7.75) on the continental shelf of western North America from Queen Charlotte Sound, Canada, to San Gregorio Baja California Sur, Mexico. On transect line 5, the corrosive water reaches all the way to the surface in the inshore waters near the coast. The black dots represent station locations.

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centration is calculated from the dissolved inorganic carbon (DIC) and total alkalinity (TA) measurements (II). In regions where  $\Omega_{\rm arag}$  or  $\Omega_{\rm cal}$  is > 1.0, the formation of shells and skeletons is favored. Below a value of 1.0, the water is corrosive and dissolution of pure aragonite and unprotected aragonite shells will begin to occur (I2). Recent studies have shown that in many regions of the ocean, the aragonite saturation horizon shoaled as much as 40 to 200 m as a direct consequence of the uptake of anthropogenic CO<sub>2</sub> (I3, I3, I4). It is shallowest in the northeastern Pacific Ocean, only 100 to 300 m from the ocean

surface, allowing for the transport of undersaturated waters onto the continental shelf during periods of upwelling.

In May and June 2007, we conducted the North American Carbon Program (NACP) West Coast Cruise on the Research Ship *Wecoma* along the continental shelf of western North America, completing a series of 13 cross-shelf transects from Queen Charlotte Sound, Canada, to San Gregorio Baja California Sur, Mexico (Fig. 1). Full water column conductivity-temperature-depth rosette stations were occupied at specified locations along each transect (Fig. 1). Water samples were

15 14 13 12 25.2 Depth (m) 25.6 Colors: 10 9 8 7 6 5 4 26 Temp (°C) 26.4 Contours: 200 Potential 26.6 26.6 Density В 0 3.5 2.2 3.0 Depth (m) 2.5 -1.2 2.0 1.5 Aragonite 1.0 200 Saturation 0.5 State 0.0 C 8.5 8.1 1.8 8.3 Depth (m) 8.1 7.9 7.7 200 7.5 рΗ 7.3 D 0 2350 2000 2000 2250 **Depth (m)** 2150 2050 Dissolved Inorganic 2220 1950 200 Carbon 2240 (µmol kg-1) 1850 Ε 0 1450 350 1300 400 1150 Depth (m) 500 1000 700 1200 850 900 700 1000 550 pCO<sub>2</sub> 200 400 (µatm) 1100 250

**Fig. 2.** Vertical sections of (**A**) temperature, (**B**) aragonite saturation, (**C**) pH, (**D**) DIC, and (**E**)  $pCO_2$  on transect line 5 off Pt. St. George, California. The potential density surfaces are superimposed on the temperature section. The 26.2 potential density surface delineates the location of the first instance in which the undersaturated water is upwelled from depths of 150 to 200 m onto the shelf and outcropping at the surface near the coast. The red dots represent sample locations.

125.5°W

126°W

125°W

collected in modified Niskin-type bottles and analyzed for DIC, TA, oxygen, nutrients, and dissolved and particulate organic carbon. Aragonite and calcite saturation, seawater pH (pH<sub>SW</sub>), and partial pressure of CO<sub>2</sub> (*p*CO<sub>2</sub>) were calculated from the DIC and TA data (11).

The central and southern coastal region off western North America is strongly influenced by seasonal upwelling, which typically begins in early spring when the Aleutian low-pressure system moves to the northwest and the Pacific High moves northward, resulting in a strengthening of the northwesterly winds (13, 14). These winds drive net surface-water Ekman transport offshore, which induces the upwelling of CO<sub>2</sub>-rich, intermediate-depth (100 to 200 m) offshore waters onto the continental shelf. The upwelling lasts until late summer or fall, when winter storms return.

During the cruise, various stages and strengths of upwelling were observed from line 2 off central Vancouver Island to line 11 off Baja California, Mexico. We observed recent upwelling on lines 5 and 6 near the Oregon-California border. Coincident with the upwelled waters, we found evidence for undersaturated, low-pH seawater in the bottom waters as depicted by  $\Omega_{arag}$  values < 1.0 and pH values < 7.75. The corrosive waters reached mid-shelf depths of ~40 to 120 m along lines 2 to 4 and lines 7 to 13 (Fig. 1). In the region of the strongest upwelling (line 5), the isolines of  $\Omega_{\text{arag}} = 1.0$ , DIC = 2190, and pH = 7.75 closely followed the 26.2 potential density surface (Fig. 2). This density surface shoaled from a depth of ~150 m in the offshore waters and breached the surface over the shelf near the 100-m bottom contour, ~40 km from the coast. This shoaling of the density surfaces and CO<sub>2</sub>-rich waters as one approaches land is typical of strong coastal upwelling conditions (15-18). The surface-water pCO<sub>2</sub> on the 26.2 potential density surface was about 850 uatm near the shelfbreak and higher inshore (Fig. 2), possibly enhanced by respiration processes on the shelf (17). These results indicate that the upwelling process caused the entire water column shoreward of the 50-m bottom contour to become undersaturated with respect to aragonite, a condition that was not predicted to occur in openocean surface waters until 2050 (5). On line 6, the next transect south, the undersaturated water was close to the surface at ~22 km from the coast. The lowest  $\Omega_{arag}$  values (<0.60) observed in the nearbottom waters of the continental shelf corresponded with pH values close to 7.6. Because the calcite saturation horizon is located between 225 and 400 m in this part of the northeastern Pacific (19), it is still too deep to shoal onto the continental shelf. Nevertheless, the calcite saturations values drop in the core of the upwelled water ( $\Omega_{cal} < 1.3$ ).

As noted, the North Pacific aragonite saturation horizons are among the shallowest in the global ocean (3). The uptake of anthropogenic CO<sub>2</sub> has caused these horizons to shoal by 50 to 100 m since preindustrial times so that they are within the density layers that are currently being upwelled along the west coast of North America.

Although much of the corrosive character of these waters is the natural result of respiration processes at intermediate depths below the euphotic zone, this region continues to accumulate more anthropogenic  $\mathrm{CO}_2$  and, therefore, the upwelling processes will expose coastal organisms living in the water column or at the sea floor to less saturated waters, exacerbating the biological impacts of ocean acidification.

On the basis of our observed O2 values and estimated O<sub>2</sub> consumption rates on the same density surfaces (18-20), the upwelled water off northern California (line 5) was last at the surface about 50 years ago, when atmospheric CO<sub>2</sub> was about 65 ppm lower than it is today. The open-ocean anthropogenic CO2 distributions in the Pacific have been estimated previously (4, 19, 21). By determining the density dependence of anthropogenic CO<sub>2</sub> distributions in the eastern-most North Pacific stations of the Sabine et al. (21) data set, we estimate that these upwelled waters contain  $\sim 31 \pm 4 \,\mu\text{mol kg}^$ anthropogenic CO<sub>2</sub> (fig. S2). Removing this signal from the DIC increases the aragonite saturation state of the waters by about 0.2 units. Thus, without the anthropogenic signal, the equilibrium aragonite saturation level ( $\Omega_{arag} = 1$ ) would be deeper by about 50 m across the shelf, and no undersaturated waters would reach the surface. Water already in transit to upwelling centers carries increasing anthropogenic CO<sub>2</sub> and more corrosive conditions to the coastal oceans of the future. Thus, the undersaturated waters, which were mostly a problem for benthic communities in the deeper waters near the shelf break in the preindustrial era, have shoaled closer to the surface and near the coast because of the additional inputs of anthropogenic CO<sub>2</sub>.

These observations clearly show that seasonal upwelling processes enhance the advancement of the corrosive deep water into broad regions of the North American western continental shelf. Because the region experiences seasonal periods of enhanced aragonite undersaturation, it is important to understand how the indigenous organisms deal with this exposure and whether future increases in the range and intensity of the corrosiveness will affect their survivorship. Presently, little is known about how this intermittent exposure to corrosive water might affect the development of larval, juvenile, and adult stages of aragonitic calcifying organisms or finfish that populate the neritic and benthic environments in this region and fuel a thriving economy. Laboratory and mesocosm experiments show that these changes in saturation state may cause substantial changes in overall calcification rates for many species of marine calcifiers including corals, coccolithophores, foraminifera, and pteropods, which are a major food source for local juvenile salmon (8, 22-30). Similar decreases in calcification rates would be expected for edible mussels, clams, and oysters (22, 31). Other research indicates that many species of juvenile fish and shellfish of economic importance to coastal regions are highly sensitive to higher-than-normal CO<sub>2</sub> concentrations such that high rates of mortality are directly correlated with the higher CO2 concentrations (31, 32). Although comprehensive field studies of organisms and their response to sporadic

increases in  $\mathrm{CO}_2$  along the western North American coast are lacking, current studies suggest that further research under field conditions is warranted. Our results show that a large section of the North American continental shelf is affected by ocean acidification. Other continental shelf regions may also be affected where anthropogenic  $\mathrm{CO}_2$ -enriched water is being upwelled onto the shelf.

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#### Supporting Online Material

www.sciencemag.org/cgi/content/full/1155676/DC1 Materials and Methods Figs. S1 and S2 References

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# Regulation of Hepatic Lipogenesis by the Transcription Factor XBP1

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Dietary carbohydrates regulate hepatic lipogenesis by controlling the expression of critical enzymes in glycolytic and lipogenic pathways. We found that the transcription factor XBP1, a key regulator of the unfolded protein response, is required for the unrelated function of normal fatty acid synthesis in the liver. XBP1 protein expression in mice was elevated after feeding carbohydrates and corresponded with the induction of critical genes involved in fatty acid synthesis. Inducible, selective deletion of XBP1 in the liver resulted in marked hypocholesterolemia and hypotriglyceridemia, secondary to a decreased production of lipids from the liver. This phenotype was not accompanied by hepatic steatosis or compromise in protein secretory function. The identification of XBP1 as a regulator of lipogenesis has important implications for human dyslipidemias.

epatic lipid synthesis increases upon ingestion of excess carbohydrates, which are converted into triglyceride (TG) in the liver and transported to adipose tissue for energy storage. Dysregulation of hepatic lipid metabolism is closely related to the development of metabolic syndrome, a condition characterized by central obesity, dyslipidemia, elevated blood glucose, and hypertension (1). In mammals, hepatic lipid metabolism is controlled by transcription factors, such as liver X receptor (LXR), sterol

regulatory element-binding proteins (SREBPs), and carbohydrate response element-binding protein (ChREBP), that regulate the expression of

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