Early Pliocene increase in thermohaline overturning: 
A precondition for the development of the modern equatorial Pacific cold tongue

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[1] Unraveling the processes responsible for Earth’s climate transition from an “El Niño–like state” during the warm early Pliocene into a modern-like “La Niña–dominated state” currently challenges the scientific community. Recently, the Pliocene climate switch has been linked to oceanic thermocline shoaling at ~3 million years ago along with Earth’s final transition into a bipolar icehouse world. Here we present Pliocene proxy data and climate model results, which suggest an earlier timing of the Pliocene climate switch and a different chain of forcing mechanisms. We show that the increase in North Atlantic meridional overturning circulation between 4.8 and 4.0 million years ago, initiated by the progressive closure of the Central American Seaway, triggered overall shoaling of the tropical thermocline. This preconditioned the turnaround from a warm eastern equatorial Pacific to the modern equatorial cold tongue state about 1 million years earlier than previously assumed. Since ~3.6–3.5 million years ago, the intensification of Northern Hemisphere glaciation resulted in a strengthening of the trade winds, thereby amplifying upwelling and biogenic productivity at low latitudes.


1. Introduction

[2] Understanding the causal chain of Pliocene climate forcing may help to improve predictions of future climate change, including the ultimate role of the ocean circulation in a globally warmer world [Jansen et al., 2007]. During the Pliocene warm period from ~5.5 to ~3 Ma, mean global surface temperatures were ~3°C warmer than today, atmospheric CO2 concentrations were close to modern ones [Foster et al., 2009], the modern Northern Hemisphere ice cap was absent, and sea level was ~25 m higher than today [e.g., Raymo et al., 1996; Dowsett et al., 1999; Ravelo et al., 2004; Mudelsee and Raymo, 2005]. Growing evidence from early Pliocene paleoclimate data suggests that sea surface temperatures (SST) in the eastern equatorial Pacific (EEP) cold tongue were similar to those of the western tropical Pacific Warm Pool [Chaisson and Ravelo, 2000; Molnar and Cane, 2002; Ravelo et al., 2004; Wara et al., 2005]. Warm surface water, a deep thermocline, and low biogenic productivity characterized the low-latitude to midlatitude upwelling regions [Fedorov et al., 2006; Dekens et al., 2007].

[3] The timing and mechanisms for the development of the modern EEP cold tongue state have so far been linked to the mid-Pliocene intensification of Northern Hemisphere glaciation (NHG) [Fedorov et al., 2006]. According to a recent hypothesis [Fedorov et al., 2006], cooling of the deep ocean, associated with enhanced oceanic heat loss at high latitudes and a balanced increase in heat gain at low latitudes, caused the tropical thermocline to shoal. This thermocline shoaling has been proposed to reach a critical threshold at ~3 Ma, allowing trade winds to bring cooler waters to the surface in equatorial and coastal upwelling zones.

[4] However, there is also evidence that cooling of the upwelling regions off California [Fedorov et al., 2006; Dekens et al., 2007; Liu et al., 2008; Brierley et al., 2009], West Africa [Marlow et al., 2000] and within the EEP cold tongue [Lawrence et al., 2006; Dekens et al., 2007].
started more than 600 kyr earlier (~4.5–3.6 Ma). Here, we present new proxy data that strongly corroborate an early development of the EEP cold tongue. This timing questions the intensification of NHG as exclusive forcing mechanism. Instead, it suggests an additional link to the early Pliocene threshold in the closure history of the Central American Seaway (CAS), which was associated with an increase in the intensification of NHG as exclusive forcing mechanism. We present new Pliocene planktonic foraminiferal δ18O and Mg/Ca-derived temperatures, alkenone (U37K) SSTs, opal accumulation rates, and benthic foraminiferal δ13C from tropical eastern Pacific Ocean Drilling Program (ODP) sites 1239 (0°40′S, 82°05′W, 1414 m water depth) and 1241 (5°51′N, 86°27′W, 2027 m water depth) [Mix et al., 2003] as well as from Caribbean ODP sites 999 and 1000. Due to the lower amount of planktonic foraminifers in samples from equatorial eastern Pacific Site 1239, we picked ten to fifteen specimens of *Globigerinoides ruber* (thermocline dweller) and Globorotalia tumida (habitat near the bottom of the photic zone) from the >250 μm size fraction. For benthic δ13C measurements at Site 1000, we selected one to five tests of the epibenthic foraminifera *Cibicidoides wuellerstorfi* (>315 μm). Prior to analysis, visibly contaminated specimens were slightly crushed and ultrasonically cleaned with methanol. The excess liquid and mud were siphoned off and the samples were dried at 60°C.

### 2. Stable Isotope Measurements

[6] The samples were freeze-dried and washed through a 63 μm mesh. For δ18O measurements on planktonic foraminifera, ten to fifteen specimens of *Neogloboquadrina dutertrei* (thermocline dweller) were selected from the 315–400 μm size fraction of samples from Caribbean sites 999 and 1000. Due to the lower amount of planktonic foraminifers in samples from equatorial eastern Pacific Site 1239, we picked ten to fifteen specimens of *Globigerinoides sacculifer* (mixed layer dweller) and *Globorotalia tumida* (habitat near the bottom of the photic zone) from the >250 μm size fraction. For benthic δ13C measurements at Site 1000, we selected one to five tests of the epibenthic foraminifera *Cibicidoides wuellerstorfi* (>315 μm). Prior to analysis, visibly contaminated specimens were slightly crushed and ultrasonically cleaned with methanol. The excess liquid and mud were siphoned off and the samples were dried at 60°C.

[7] Stable isotope analyses on planktonic and epibenthic foraminifers were carried out at IFM-GEOMAR (Kiel), using either a Finnigan Delta-Plus-Advantage mass spectrometer coupled to a Finnigan Gas Bench II (with analytical precision better than ±0.07 %o for δ18O and better than ±0.05 %o for δ13C; ±1σ), or a Finnigan MAT 252 mass spectrometer with automated Kiel carbonate preparation device (with analytical precision better than ±0.07 %o for δ18O and ±0.04 %o for δ13C; ±1σ). Both machines were intercalibrated using a house standard, which was calibrated to the National Bureau of Standards NBS-19. The ratios of δ18O/δ16O and δ13C/δ12C are reported with reference to the Pee Dee Belemnite (PDB) standard.
2.3. Mg/Ca Analyses

For Mg/Ca measurements, 20–25 specimens of *N. dutertrei* (sites 999, 1000) and *G. tumida* (Site 1239) were picked from the same size fraction as used for stable isotope analyses. Specimens visibly contaminated by ferromanganese oxides were not selected for analysis. After gentle crushing, the samples were cleaned according to the cleaning protocol for Mg/Ca of Barker et al. [2003]. To remove clays, the samples were rinsed four to six times with distilled deionized water and twice with methanol (suprapure) with ultrasonic cleaning steps (2 to 3 min) after each rinse. Subsequently, samples were treated with a hot (97°C) oxidizing 1% NaOH/H2O2 solution (10 ml 0.1 N NaOH (analytical grade); 100 μl 30% H2O2 (suprapure)) for 10 min to remove organic matter. Every 2.5 min, the vials were rapped on the bench top to release any gaseous buildup. After 5 min, the samples were placed in an ultrasonic bath for a few seconds to maintain contact between reagent and sample. This treatment was repeated after refreshment of the oxidizing solution. The remaining oxidizing solution was removed during three rinsing steps with distilled deionized water. After transferring the samples into clean vials, a weak acid leach with 250 μl 0.001 M HNO3 (subboiled distilled) was applied with 30 s ultrasonic treatment and subsequent two rinses with distilled deionized water. After cleaning, the samples were dissolved in 0.075 M nitric acid (HNO3) (subboiled distilled) and diluted several times, until all samples obtained calcium concentrations in the range of 30–70 ppm.

Analyses were performed on an ICP-AES (ISA Jobin Yvon–Spex Instruments S.A. GmbH) at IFM-GEOMAR (Kiel) or on a simultaneous, radially viewing ICP-OES (Ciros CCD SOP, Spectro A. I., Germany) at the Institute of Geosciences (Kiel University, Germany). The long-term precision of the Mg/Ca analyses estimated from an internal laboratory standard was 0.1%. Replicate analyses on the same samples, which were cleaned and analyzed during different sessions, showed a standard deviation of 0.09 mmol/mol, introducing a temperature error of about 0.5°C.

2.4. Calculation of Mg/Ca-Derived Paleotemperatures

For this study, *G. sacculifer* Mg/Ca temperatures were calculated using the Dekens et al. [2002] Mg/Ca temperature calibration for *G. sacculifer* with core-depth-based dissolution corrections for the Pacific (Site 1241) and for the Atlantic (sites 999, 1000). *N. dutertrei* Mg/Ca ratios from Caribbean sites 999 and 1000 were converted into temperature using the Dekens et al. [2002] Mg/Ca temperature calibration for *N. dutertrei* with a core-depth-based dissolution correction for the Atlantic. We used the Anand et al. [2003] multispecies Mg/Ca temperature calibration to calculate *G. tumida* Mg/Ca paleotemperatures at Pacific sites 1239 and 1241 and we included the Dekens et al. [2002] core-depth-based Pacific dissolution correction for *G. sacculifer*, because species-specific dissolution corrections for *G. tumida* are not available for the Pacific.

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\text{Mg/Ca} = 0.38 \exp(0.09[T - 0.36(\text{core depth in km}) - 2.0°C])
\]

[11] Yet we are aware that Mg/Ca *G. tumida* might have a different sensitivity to carbonate dissolution than Mg/Ca *G. sacculifer*, which would in turn affect the resulting absolute *G. tumida* Mg/Ca temperature estimates. Absolute Mg/Ca temperature estimates of course depend on the selection of Mg/Ca temperature calibrations used for the calculations, but the relative changes in vertical temperature gradients are robust regardless of which calibration is used. Our conclusions, which are mainly based on the evolutionary long-term changes of vertical temperature gradients between the mixed layer and the bottom of the photic zone, would therefore not be affected by the application of other Mg/Ca temperature calibrations.

2.5. Calculation of Opal Accumulation Rates

To estimate opal accumulation rates at Site 1239, we quantified the concentrations of biogenic silica (opal) for 15 samples by using the automated leaching method according to Müller and Schneider [1993]. The opaline material was extracted from the dry and ground bulk sediment by sodium hydroxide at ∼85°C for ∼45 min. The leaching solution was continuously analyzed for dissolved silicon by molybdate blue spectrophotometry. The DeMaster [1981] mineral correction was consequently applied. The excellent linear correlation of opal concentrations with shipboard grape density data [Mix et al., 2003] (R = 0.8) allowed for the calculation of a high-resolution opal percentage record. Mass accumulation rates (g cm⁻² kyr⁻¹) were then calculated using the sedimentation rate (in cm kyr⁻¹) between age control points [Tiedemann et al., 2007], dry bulk density data calculated from grape density and shipboard physical properties data [Mix et al., 2003], and the opal concentrations (%).

2.6. Alkenone Analysis

Samples of 2 g freeze-dried and homogenized sediment were mixed with an internal standard and ultrasonically extracted for 3 min (UP200H ultrasonication disrupter probe; S3 micropoint, amplitude 0.5, pulse 0.5), using successively less polar mixtures of methanol and methylene chloride (CH3OH, CH3OH/CH2Cl2 1:1, CH2Cl2). After centrifuging, the supernatants were combined, desalted with deionized water, dried with Na2SO4 and rotary evaporated to complete dryness. The residues were dissolved in CH3Cl2 and additionally purified using a silica cartridge (Varian Bond Elut; 1CC/100 mg). To eliminate interference with wax esters, the clean extracts were hydrolyzed with 0.1 N KOH in Methanol (90/10 CH3OH/H2O) at 80°C for 2 h, and the neutral fraction containing the alkenones was obtained by partitioning into hexane. Finally, the extracts were concentrated under N2 and taken up in 25 μl of the 1:1 CH3OH/CH2Cl2 mixture. Gas chromatography was performed using a HP5890 series II gas chromatograph equipped with a split/spillless injector, a 60 m × 0.32 mm × 0.1 μm nonpolar fused silica capillary column DB-S5MS and flame ionization detector. An aliquot of 3 μl was injected in split mode (one tenth) with helium as the carrier gas. The oven temperature was programmed from 50°C to 250°C at 25°C/min, 250°C to 290°C at 1°C/min and a final heating from 290°C to 310°C at 30°C/min. Compounds were quantified using octacosane acid methyl ester as an internal standard and the relative response
of the C38 n-alkane. The ketone unsaturation index $U_{37}^K$ was converted to temperature according to Conte et al. [2006]:

$$T(°C) = -0.957 + 54.3\left(U_{37}^K\right) - 52.9\left(U_{37}^K\right)^2 + 28.3\left(U_{37}^K\right)^3.$$  

(2)

[14] We applied the nonlinear $U_{37}^K$/temperature calibration of Conte et al. [2006] because it is based on alkenones from surface water particulates collected in the world ocean (i.e., the natural environment of the alkenone producers) and associated temperatures measured at sampling depth. Moreover, this calibration accounts for the asymptotic behavior of $U_{37}^K$ at the warmest SSTs. The analytical precision ($±1σ$) based on duplicates and multiple extractions of a sediment sample used as an internal laboratory reference sample was better than 0.003 $U_{37}^K$ units or 0.1°C.

2.7. Climate Model Experiments

[15] For our sensitivity experiments, we used the global atmosphere-ocean model ECBILT-CLIO version 3. The coupled model derives from the atmosphere model ECBILT [Opsteegh et al., 1998] and the ocean/sea ice model CLIO [Goosse and Fichefet, 1999]. The atmospheric component solves the quasi-geostrophic equations and ageostrophic correction terms in T21 resolution using three layers. The primitive equation, free surface ocean component has a horizontal resolution of 3 degrees and 20 levels in the vertical. It is coupled to a thermodynamic-ageostrophic sea ice model with viscous plastic rheology. There is no local flux correction in ECBILT-CLIO. However, precipitation over the Atlantic and Arctic basins is reduced by 8.5% and 25%, respectively, and homogeneously redistributed over the North Pacific. The implementation of this regional flux adjustment considerably improves the simulation of the present-day climate and produces a realistic AMOC. Its reliability in the simulation of other climates than the present one, however, is elusive. For the sensitivity studies presented in this paper, the potential errors in surface hydrography and AMOC that are induced by the flux adjustment are acceptable since we are interested in the qualitative relationship between the AMOC and the thermocline, rather than in precise numbers. More information about the model and a complete list of references is available at http://www.knmi.nl/onderzk/CKO/ecbilt-papers.html.

[16] Beside a 5000 years control run (experiment CTL), simulating the present-day (preindustrial) climate, we conducted an experiment with open Central American Seaway (experiment CAS) and a freshwater hosing experiment (experiment OFF). These experiments were designed to elucidate the relationship between AMOC and tropical TCD. The freshwater hosing is a convenient method to shut off the AMOC, providing an estimate for the maximum response of tropical TCD to pure AMOC slowing. We note that we do not imply corresponding meltwater events in the early Pliocene.

[17] In experiment CAS, the Panama Seaway has a sill depth of 700 m and is defined on three velocity grid points, corresponding to a width of approximately 1000 km. All other boundary conditions are the same as in the control run. Therefore, experiment CAS should be viewed as a sensitivity study to elucidate tropical thermocline dynamics rather than a simulation of early Pliocene climate. In experiment OFF Panama is closed, but NADW formation is completely shut off by an anomalous freshwater input to the North Atlantic (0.5 Sv between 50°N and 70°N). In both experiments CAS and OFF, the model was integrated another 2250 years to reach a new equilibrium, starting from the control run’s final state as initial condition. Model results presented in this paper are annual averages determined from the last 50 years of each experiment.

[18] Generally, the thermocline is defined as the depth at which the vertical temperature gradient is at a maximum. In a numerical ocean model such as CLIO, the search for this maximum would always result in thermocline depths that correspond exactly to the depths of grid points (in ECBILT-CLIO these are located at 5 m, 16 m, 29 m, 45 m, 65 m, 90 m, 122 m, etc.). Hence, changes that are smaller than the vertical grid spacing could not be observed when using the maximum temperature gradient for the calculation of TCD. Therefore, alternative indicators for TCD are generally used in numerical models. As an indicator for tropical TCD, we use the depth of the 20°C isotherm (the depth of an isotherm is not bound to the grid spacing; an isotherm can well reside between two grid points in the vertical and its depth can easily be found by linear interpolation of the gridded temperature field). In both observational data and the ECBILT-CLIO climate model, the depth of the maximum temperature gradient is almost identical to the depth of the 20°C isotherm in the central (~150 m) and eastern (less than 50 m on average) equatorial Pacific.

3. Results

3.1. Equatorial Eastern Pacific Site 1239

3.1.1. Planktonic $δ^{18}O$ and Mg/Ca Paleotemperature Records

[19] The $δ^{18}O$ record of the mixed layer dweller $G. sacculifer$ as well as $δ^{18}O$ and Mg/Ca temperature records of the deep dweller $G. tumida$ (habitat close to the bottom of the photic zone [Ravelo and Fairbanks, 1992; Ravelo and Andreasen, 1999]) serve to reconstruct Pliocene changes in upper ocean stratification at equatorial eastern Pacific Site 1239 (Figure 2a). Small (large) surface/subsurface temperature differences, and hence small (large) $δ^{18}O$ differences between shallow- and deep-dwelling planktonic foraminifera indicate a deep (shallow) thermocline [e.g., Ravelo and Fairbanks, 1992; Ravelo and Andreasen, 1999; Steph et al., 2009].

[20] At Site 1239, $δ^{18}O$ values of the mixed layer dweller $G. sacculifer$ remained relatively constant between 5.0 and 3.9 Ma (average $δ^{18}O$ of −1.1‰). From 3.9 to 2.7 Ma, the $δ^{18}O$ values of $G. sacculifer$ increased by ~0.4‰ (average $δ^{18}O$ of −0.7‰ between 2.7 and 2.9 Ma) with a major step around −3.7 Ma (Figure 2a). The Mg/Ca temperature and $δ^{18}O$ records of the deep dweller $G. tumida$, however, suggest larger temperature changes at the bottom of the photic zone. Average $G. tumida$ $δ^{18}O$ values increased by
Figure 2. New proxy data presented in this study, spanning the time interval from ∼5.6 Ma to 2.1 Ma. (a) *G. sacculifer* δ¹⁸O data (red line), *G. tumida* δ¹⁸O data (light blue line) and Mg/Ca temperature data (dark blue) from equatorial eastern Pacific ODP Site 1239. (b) Biogenic opal accumulation rates, Site 1239 (green line). (c) U⁴³⁷ of sea surface temperature data from tropical eastern Pacific Site 1241 (red). (d) Epibenthic δ¹³C record from Caribbean Site 1000 (*C. wuellerstorfi*, black line). (e) The δ¹⁸O records of the thermocline dweller *N. dutertrei* (dextral variety) from Caribbean sites 999 (light blue) and 1000 (orange) and Mg/Ca temperature records of *N. dutertrei* (dextral variety) from sites 999 (dark blue) and 1000 (red).
~0.5\% between 4.8 and 4.0 Ma, and by an additional
~0.3\% between 3.8 and 3 Ma (Figure 2a). The low-resolution
G. tumida Mg/Ca temperature record at Site 1239 shows a
similar pattern; temperatures decrease by ~5°C during the
eyear Pliocene, with a major step between ~4.8 and ~4.0 Ma
(Figure 2a). This subsurface cooling and the increasing δ18O
gradient between the mixed layer and the bottom of the photic
zone point to a TCD decrease at Site 1239 during the early
Pliocene. Between 4.0 and 2.7 Ma, G. tumida Mg/Ca tem-
peratures remained relatively stable, except for a pronounced
cool interval between 3.3 and 3.0 Ma (Figure 2a).

3.1.2. Biogenic Opal Accumulation Rates
[21] We use biogenic opal accumulation rates calculated
from equatorial eastern Pacific Site 1239 shipboard grape
density data [Mix et al., 2003] as an estimate for changes in
diatom productivity (Figure 2b). Between 5.0 and 3.6 Ma,
opal accumulation rates were low (on average ~0.4 g cm⁻²
kyr⁻¹) and the variability was relatively small, except for a
short interval with higher opal accumulation rates (up to
0.7 g cm⁻² kyr⁻¹) around 4.0 Ma. Between 3.6 and 3.5 Ma,
opal accumulation rates at Site 1239 doubled rapidly to
average values of ~0.8 g cm⁻² kyr⁻¹ (reaching maximum
values of 1.05 g cm⁻² kyr⁻¹), suggesting a distinct increase of
diatom productivity at Site 1239 after 3.6–3.5 Ma (Figure 2b).

3.2. Tropical Eastern Pacific Site 1241
[22] We measured a low-resolution U²⁰ record spanning
the time interval from 4.0 to 2.5 Ma in order to get hold of
SST changes in the upper 10 m of the water column, the
preferred habitat of alkene-producing coccolithophores.
The Site 1241 U²⁰ temperatures vary between 27.4°C and
28.3°C and show no long term trend (Figure 2c), suggesting
that Pliocene SSTs in the uppermost water column were
relatively constant in time and only slightly warmer than
today (modern SSTs are ~27.0°C at the 4 Ma paleolocation
and 27.4°C at the 2 Ma paleolocation of Site 1241).

3.3. Caribbean Sites 999 and 1000
3.3.1. Site 1000 Benthic δ¹³C Record (C. wuellerstorfi)
[23] The δ¹³C record of the epibenthic foraminifer C.
wuellerstorfi serves as a proxy for changes in deep water
ventilation [e.g., Zahn et al., 1986; McCorkle and Keigwin,
1994], because the δ¹³C of seawater is closely linked to
nutrient and oxygen contents, whereas higher δ¹³C values
indicate lower nutrient contents and better ventilation
[Kroopnick, 1985]. Caribbean Site 1000 (916 m water
depth) provides information about ventilation changes at the
Atlantic intermediate water level. Benthic δ¹³C at Site 1000
deprecated during the earliest Pliocene, reaching the lowest
level between ~5.2 Ma and 4.9 Ma (average δ¹³C = 0.6 %o)
(Figure 2d). Between ~4.9 and 3.6 Ma, Site 1000 benthic
δ¹³C increased to average values of ~1.1 %o. These high δ¹³C
values persisted until 2.2 Ma. The overall early Pliocene δ¹³C
trend suggests an increase in intermediate water ventilation
(Figure 2d).

3.3.2. N. dutertrei δ¹⁸O and Mg/Ca Paleotemperature
Records
[24] We use δ¹⁸O and Mg/Ca temperature records mea-
sured on the thermoline dweller N. dutertrei in order to
reconstruct Pliocene Caribbean subsurface temperature
changes, because the deep dweller G. tumida is absent in the
Pliocene sections of sites 999 and 1000. The subsurface
temperature development was very similar at both Caribbean
sites (Figure 2e). Between 4.8 and 4.3 Ma, average δ¹⁸O,
dutertrei decreased by more than 1 %o, whereas the N. dutertrei
Mg/Ca temperature records indicate a subsurface thermal temperature increase of ~4°C. Between 4.3 and 4.0 Ma,
δ¹⁸O, dutertrei increased while N. dutertrei Mg/Ca tem-
peratures stayed stable, hinting to an increase in Caribbean
subsurface salinity (Figure 2e). After 4.0 Ma, small long-
term variations in the δ¹⁸O and Mg/Ca temperature records of
N. dutertrei indicate no major change in thermocline
temperature and/or salinity at Caribbean Site 999.

3.4. Model Results
[25] Introducing a Panama Seaway in experiment CAS
results in a mean total volume transport of 14 Sv (1 Sv =
10⁶ m³ s⁻¹) from the Pacific into the Atlantic over the entire
depth (700 m) of the strait. The inflow of relatively fresh
Pacific surface water masses into the Atlantic reduces the
salinity contrast between the two oceans and, as a result, the
AMOC decreases. This is a robust result which has already
been found in previous modeling studies and which is
effectively independent of other boundary conditions [cf.
Maier-Reimer et al., 1999; Klocker et al., 2005; Lunt et al.,
2008]. In our model, convection in the Labrador Sea,
and hence UNADW formation, stops. The strength of the
AMOC (measured by the net export of NADW to the
Southern Ocean) is 15 Sv in the control run and 9 Sv in
experiment CAS [cf. Franke and Schulz, 2004]. Figure 3a
illustrates global changes in the depth of the 20°C isotherm
(which is a good indicator for tropical TCD; see above) in
response to the closure of the CAS. The pattern indicates
an overall decrease in tropical TCD, except for the Caribbean,
where the thermocline deepens (Figure 3a). Except for
the Caribbean, the global pattern of TCD change between
experiments CTL and OFF provides a very similar picture,
albeit with a larger magnitude of thermocline shoaling in
most areas of the world ocean as compared to experiment
CAS (Figure 3b).

4. Discussion
4.1. Changes in Tropical Eastern Pacific Thermocline
Depth
[26] We use new and previously published Mg/Ca tem-
perature records and δ¹⁸O values of the shallow-dwelling
planktonic foraminifer G. sacculifer and of the deep dweller
G. tumida to assess Pliocene variations in tropical eastern
Pacific TCD (Figure 4). ODP Site 1241 is located at ~6°N
close to the present-day Intertropical Convergence Zone
(ITCZ) at the southern edge of the East Pacific Warm Pool,
where upwelling is absent and tropical wind stress conver-
gence is largest (doldrums; Figure 1). It has therefore
an ideal position to monitor oceanic adjustments in TCD.
Pliocene alkene-derived SSTs at Site 1241 were high
and relatively constant (~28°C; Figure 4a) and temperature
fluctuations derived from Mg/Ca, sacculifer were also small
(~2°C) [Groeneveld et al., 2006] (Figure 4a). Pliocene
Mg/Ca$_{G.\text{sacculifer}}$ paleotemperature estimates at Site 1241 are on average $\sim 1^\circ$C lower than alkenone-derived temperatures if using the Dekens et al. [2002] G. sacculifer calibration with Pacific dissolution correction factors for Mg/Ca temperature calculations. This relatively small temperature offset may either be related to differences in seasonality or to the fact that U37K$'\text{provides }$SST's within the uppermost water column ($\sim$0–10 m water depth) while Mg/Ca$_{G.\text{sacculifer}}$ displays mixed layer temperatures (the modern average calcification depth of G. sacculifer in the Panama Basin is $\sim$30–50 m [e.g., Fairbanks et al., 1982]). Today, the temperature difference between 0 m and 30 m water depth at the 4–2 Ma paleolocations of Site 1241 amounts to $\sim$1.5°C.

[27] In contrast, large temperature changes occurred at the bottom of the photic zone (Figure 4a). At East Pacific Warm Pool Site 1241, the temperature record of G. tumida reveals a long-term decrease of 6°C during the early Pliocene with a major step between $\sim$4.8 and $\sim$4.0 Ma [Steph et al., 2006b]. The corresponding increase in the temperature gradient between the mixed layer and the bottom of the photic zone has been interpreted as a decrease in TCD [Steph et al., 2006b]. The thermocline at Site 1241 remained shallow between $\sim$4.0 and $\sim$3.0 Ma. The subsurface temperature changes monitored by Mg/Ca$_{G.\text{tumida}}$ at equatorial upwelling Site 1239 off Ecuador (Figures 1 and 4a) are very similar to those observed at Site 1241, and also point to thermocline shoaling during the early Pliocene. Since Pliocene changes in Mg/Ca temperatures and $\delta^{18}$O recorded by G. tumida show approximately the same pattern at sites 1241 and 1239 (Figure 4a), we are confident that $\delta^{18}$O$_{G.\text{tumida}}$ predominantly reflects temperature changes at the bottom of the photic zone (although biased by a certain amount of salinity change [Steph et al., 2006b]). The fact that the G. tumida $\delta^{18}$O records from tropical eastern Pacific sites 847 [Chaisson and Ravelo, 2000; Wara et al., 2005] and 851 [Cannariato and Ravelo, 1997] bear a striking similarity to those from sites 1239 and 1241 suggests that the early Pliocene TCD decrease in the tropical eastern Pacific between 4.8 and 4.0 Ma was a large-scale phenomenon that spread across tropical oceanic fronts (Figures 1 and 4b). The temperature increase at the bottom of the photic zone after $\sim$3.0 Ma observed at doldrum Site 1241 (derived from Mg/Ca$_{G.\text{tumida}}$) points to thermocline deepening outside the EEP cold tongue region (Figure 4a), and does hence not support the hypothesis of general thermocline shoaling around 3 Ma in response to global cooling [Fedorov et al., 2006].

**Figure 3.** Differences in the depth of the 20°C isotherm (annual mean) between the present-day control run CTL and (a) experiment CAS with open Central American Seaway and (b) experiment OFF with NADW formation shut off completely by a strong anomalous freshwater input to the North Atlantic.
4.2. Pliocene Changes in AMOC

The timing of the observed regional TCD changes in the eastern tropical Pacific suggests a close link to changes in AMOC, as indicated by a comparison of benthic δ¹³C data from tropical western Atlantic Site 925 [Bickert et al., 1997; Tiedemann and Franz, 1997; Shackleton and Hall, 1997] and from Caribbean Site 1000. Specifically, the shoaling of the thermocline between 4.8 and 4.0 Ma was associated with AMOC strengthening, whereas the observed thermocline deepening at Site 1241 after 3 Ma was associated with a decrease in AMOC (Figure 5).

Caribbean Site 1000, located at 916 m water depth, provides a window into the tropical Atlantic intermediate water level. Tropical Atlantic Site 925, located on the Ceará Rise at 3041 m water depth, is situated within the modern core depth of Lower North Atlantic Deep Water (LNADW) (Figure 1). The comparison of changes in δ¹³C at sites 925 and 1000 provides information about the relative proportion of northern component (high-δ¹³C) and southern component (low-δ¹³C) water and the interplay between Upper North Atlantic Deep Water (UNADW) and LNADW formation (Figure 5). Approximations for Pliocene δ¹³C end-member values indicative of Southern Ocean intermediate water (Site 1236; southeast Pacific, 1323 m water depth [Tiedemann et al., 2007]) and Southern Ocean deep water (Site 1237; southeast Pacific, 3212 m water depth [Tiedemann et al., 2007]) are provided in Figure 5 (see arrows). Unfortunately, no Pliocene end-member values are available for North
Figure 5. (top) *G. tumida* Mg/Ca temperature record from eastern Pacific Warm Pool Site 1241 (red line with dots) showing Pliocene changes in tropical eastern Pacific thermocline depth. (middle) $\Delta^{13}C$ between Site 1000 (this study) and Site 925 [Bickert et al., 1997; Tiedemann and Franz, 1997; Shackleton and Hall, 1997]. Before calculating the $\delta^{13}C$ difference between sites, both $\delta^{13}C$ records were evenly sampled at 2 kyr steps and smoothed using a 10-point running mean. This plot serves to clarify the timing of changes in the $\delta^{13}C$ difference between sites 1000 and 925 as mentioned in the text. Prior to 4.8 Ma, values < 0 suggest a large component of southern sourced water masses at the UNADW level (since intermediate water records from Caribbean Site 1000 and southeast Pacific Site 1236 [Tiedemann et al., 2007] have similar $\delta^{13}C$ values) and weak AMOC. The decrease in $\Delta^{13}C$ from ~4.8 Ma to ~4.0 Ma is marked by an increase in northern component water at the UNADW level and indicates AMOC strengthening. $\Delta^{13}C$ values around 0 between 4 Ma and 3 Ma indicate a similar degree of ventilation at the UNADW and LNADW level. As $\delta^{13}C$ values at both sites are high during this interval, this points to enhanced AMOC. $\Delta^{13}C$ values > 0 after ~3 Ma result from decreasing $\delta^{13}C$ values at LNADW Site 925. This indicates a larger component of southern sourced water masses at the LNADW level and reflects AMOC weakening, especially during harsh climate episodes. (bottom) Comparison of benthic $\delta^{13}C$ records from Caribbean Site 1000 (red, this study) and western Atlantic Site 925 (blue) [Bickert et al., 1997; Tiedemann and Franz, 1997; Shackleton and Hall, 1997]. Arrows approximate average Pliocene $\delta^{13}C$ end-members of southern sourced water masses (orange arrow is for Site 1236, 21°22’S, 81°26’W, 1323 m water depth, Southern Ocean intermediate water [Tiedemann et al., 2007]; gray arrow is for Site 1237, 16°00’S, 76°23’W, 3212 m water depth, Southern Ocean circumpolar deep water [Tiedemann et al., 2007]).
Atlantic intermediate water masses. However, $\delta^{13}$C values at Site 1000 higher than $\sim$0.5% (average Pliocene $\delta^{13}$C value at Southern Ocean intermediate water Site 1236) can only be explained by admixture of northern sourced (comparatively high $\delta^{13}$C) intermediate water masses. At the core depth of LNADW (Site 925), $\delta^{13}$C values between 5.5 and 3.0 Ma were always too high to be explained by a significant contribution of southern sourced water masses. This suggests that the $\delta^{13}$C contrast between northern and southern end-members was strong and relatively constant during the early to middle Pliocene.

[30] Our findings generally support previous studies, which documented that changes in AMOC were linked to a threshold in the closure history of the CAS during the early Pliocene and to the mid-Pliocene intensification of NHG [e.g., Keigwin, 1978; Oppo et al., 1995; Haug and Tiedemann, 1998; Haug et al., 2001]. Between 5.5 and 3.2 Ma, average $\delta^{13}$C values at Site 925 were relatively high ($\sim$1%), indicating the continuous presence of well-ventilated LNADW (Figure 5). At Site 1000, however, $\delta^{13}$C-depleted Southern Ocean intermediate water masses dominated before $\sim$4.9–4.8 Ma. After 4.8 Ma, the $\delta^{13}$C difference between sites 925 and 1000 started to decrease until benthic $\delta^{13}$C at Site 1000 reached values identical to those at Site 925 by $\sim$4.0 Ma (Figure 5). This points to a first maximum in UNADW production in the Labrador Sea. The long-term $\delta^{13}$C increase at UNADW Site 1000 was paralleled by thermocline shoaling in the tropical eastern Pacific. The intensification of UNADW formation has been attributed to enhanced salt transfer into the high northern latitudes via the Gulf Stream in response to shoaling of the CAS [e.g., Keigwin, 1978; Haug and Tiedemann, 1998].

[31] By comparing early Pliocene changes in benthic $\delta^{13}$C at sites 1000 and 925, we obtain basically the same results as Haug and Tiedemann [1998] and Haug et al. [2001], who used benthic $\delta^{13}$C records from Caribbean Site 999 (2928 m water depth), east Atlantic ODP Site 659 (3070 m water depth) and ODP Site 849 (deep Pacific end-member). It was recently questioned, however, whether benthic $\delta^{13}$C changes at Site 999 (which is located $\sim$1000 m below the Atlantic/Caribbean sill depth) indeed reflect Pliocene changes in Atlantic intermediate water circulation [Molnar, 2008]. The significant advantage of our new $\delta^{13}$C record from Site 1000 (916 m water depth) is that it monitors changes in water mass characteristics well above the Atlantic/Caribbean sill depth ($\sim$1900 m), where the exchange between Atlantic and Caribbean intermediate waters was unhampered. Although the long-term trend of $\delta^{13}$C changes is similar at sites 999 and 1000, absolute $\delta^{13}$C values at Site 1000 are significantly higher than those from Site 999 [Haug and Tiedemann, 1998]. This demonstrates the comparatively stronger influence of UNADW at the shallower Site 1000 (with $\delta^{13}$C values of up to 1.1‰; Figure 5). Such high $\delta^{13}$C values are not observed in Southern Ocean sourced deep and intermediate water masses [Tiedemann et al., 2007].

[32] High and nearly identical benthic $\delta^{13}$C values at sites 1000 (UNADW) and 925 (LNADW) persisted between $\sim$4.0 and 3.0 Ma (Figure 5). This corroborates the general consensus of a “superconveyor” with strong UNADW and LNADW formation [e.g., Raymo et al., 1996; Ravelo and Andreasen, 2000] and coincides with the establishment and persistence of a shallow thermocline in the tropical east Pacific. Along with the first appearance of larger ice sheets on northern continents, $\delta^{13}$C values started to decrease at Site 925, but remained high at Site 1000 (Figure 5). During harsh climate episodes, UNADW production stayed high (high $\delta^{13}$C values at UNADW Site 1000) whereas LNADW formation was reduced [Oppo et al., 1995], allowing poorly ventilated bottom waters from the Southern Ocean to penetrate to shallower water depths and farther to the north (low $\delta^{13}$C values at LNADW Site 925). This weakening of the AMOC was paralleled by a moderate thermocline deepening in the tropical eastern Pacific (Figure 5).

4.3. Changes in Caribbean Thermocline Depth

[33] In order to assess Pliocene variations in TCD at Caribbean sites 999 and 1000, we compare Mg/Ca temperature and $\delta^{18}$O data of the thermocline dweller N. dutertrei to previously published Mg/Ca temperature data of the mixed layer dweller G. sacculifer from Caribbean sites 999 and 1000 [Groeneveld, 2005; Groeneveld et al., 2008] (Figure 6). A recent tropical Atlantic/Caribbean core top study indicates that $\delta^{18}$O$_{N. dutertrei}$ − $\delta^{18}$O$_{G. sacculifer}$ can serve as a faithful recorder of TCD in the western Atlantic and Caribbean. Core top $\delta^{18}$O$_{N. dutertrei}$ and hence $\delta^{18}$O$_{N. dutertrei}$ − $\delta^{18}$O$_{G. sacculifer}$ decreases significantly with increasing TCD, although N. dutertrei tends to calcify deeper in the water column if the thermocline is deep [Steph et al., 2009].

[34] In this study, we use G. sacculifer Mg/Ca temperature records instead of $\delta^{18}$O$_{G. sacculifer}$ because early Pliocene mixed layer $\delta^{18}$O changes in the Caribbean were predominantly driven by salinity [e.g., Haug et al., 2001; Steph et al., 2006a]. Mixed layer temperatures at both Caribbean sites show only small variations during the early Pliocene (Figure 6), whereas the Dekens et al. [2002] dissolution-corrected Mg/Ca temperature estimates suggest $\sim$1.5°C higher temperatures at Site 1000 (on average 26.8°C) than at Site 999 (on average 25.3°C; Figure 6). Without dissolution corrections (using, i.e., the Mg/Ca temperature calibration of Anand et al. [2003]), the temperature difference between both sites would amount to $\sim$3°C. Today, temperatures at 40–80 m water depth (estimated modern calcification depth of G. sacculifer in the Caribbean [Steph et al., 2006a]) are $\sim$1°C higher at Site 1000 than at Site 999 (NODC, 2001 [Conkright et al., 2002]). Yet it is possible that this temperature difference was slightly enlarged during the early Pliocene, since Site 999 was located closer to the final openings of the CAS than Site 1000 (which was probably less affected by the comparatively cool Pacific inflow as discussed by Steph et al. [2006a] (see Figure 1)). Accordingly, mixed layer temperatures at Pacific Site 1241 and at Caribbean Site 999 should have been similar as long as Pacific surface water entered the Caribbean via the CAS (not necessarily at Site 1000, since it was located at a greater distance from the final openings of the gateway [see Steph et al., 2006a]). Indeed, the measured Mg/Ca ratios of G. sacculifer are similar at sites 999 and 1241. Using the Dekens et al. [2002] Mg/Ca temperature calibration with Pacific dissolution correction factors for Site 1241 and Atlantic dissolution correction factors for Caribbean Site
999, however, the resulting absolute mixed layer temperature estimates are generally higher at Site 1241 than at Site 999, especially between ∼4.6 and 2.5 Ma (compare Figures 4a and 6). This may either indicate that early Pliocene carbonate dissolution patterns were different from the modern ones (with relatively stronger carbonate dissolution in the Caribbean and/or enhanced carbonate preservation in the Pacific with respect to today), or that relatively cool water masses (not originating from the East Pacific Warm Pool; Site 1241) entered the Caribbean from the EEP cold tongue region. Yet this would imply that parts of the EEP cold tongue were already significantly cooler than the East Pacific Warm Pool during the early Pliocene (which is not supported by the alkenone SST record from EEP cold tongue Site 846 [Lawrence et al., 2006]).

[35] Since Pliocene mixed layer temperatures at Caribbean sites 999 and 1000 remained relatively constant through time, the observed increase in N. dutertrei Mg/Ca temperatures after ∼4.8 Ma and the resulting decrease in the temperature gradient between mixed layer (G. sacculifer) and the thermocline (N. dutertrei), as well as the contemporaneous decrease in δ¹⁸O_N. dutertrei (Figure 6) point to a deepening/warming of the Caribbean thermocline synchronous with thermocline shoaling in the tropical eastern Pacific [e.g., Cannariato and Ravelo, 1997; Steph et al., 2006b, chapter 4.1]. The timing of this event corresponds roughly to the restriction of Pacific–Caribbean surface water exchange after 4.7–4.6 Ma [e.g., Keigwin, 1982; Haug et al., 2001; Steph et al., 2006a].

4.4. Numerical Experiments

[36] In the ECBILT–CLIO experiments, the closure of the CAS induces an overall decrease in tropical TCD, except for the Caribbean, where the thermocline deepens (Figure 3a). This is consistent with our Pliocene TCD reconstructions for Caribbean sites 999 and 1000 (Figure 6), and with results from previously published modeling studies [Schneider and Schmittner, 2006]. We thus interpret the Caribbean TCD increase after ∼4.8 Ma to indicate the development of the modern Caribbean Warm Pool in response to shoaling of the CAS, when the inflow of relatively cold subsurface water from the Pacific into the Caribbean became significantly reduced [e.g., Keigwin, 1982; Haug et al., 2001; Steph et al., 2006a; Groeneveld et al., 2008]. Except for the Caribbean, the global pattern of TCD change between experiments CTL and OFF shows a very similar picture in most areas of the world ocean (Figure 3b). This result provides insight into the mechanism of thermocline shoaling: Enhanced NADW formation leads to an increase in the volume of the cold water sphere and, hence, to upward thermocline shifts in the global ocean [Huang et al., 2000; Timmermann et al., 2005]. This finding suggests that the thermocline shoaling in the tropical east Pacific during the early Pliocene was the result of a global oceanic adjustment process in response to AMOC amplification, which, in turn, was induced by substantial shoaling of the CAS. Changes in wind stress owing to AMOC intensification were unlikely to account for the thermocline shoaling in the eastern equatorial Pacific (compared to experiments CAS and OFF an eastward surface wind stress anomaly is found over the entire equatorial Pacific in CTL, which tends to deepen the tropical thermocline in the eastern equatorial Pacific; see auxiliary material).¹ However, the simulated trade wind intensification in the southeastern tropical Pacific leads to cooling via increased evaporation and Ekman pumping that may lead to local thermocline shoaling [cf. Timmermann et

¹Auxiliary materials are available in the HTML. doi:10.1029/2008PA001645.

Figure 6. Changes in Caribbean upper ocean water mass signatures at ODP sites 999 and 1000 during the time interval from 5.6 to 2.1 Ma. Mg/Ca–based mixed layer temperature records (G. sacculifer) from sites 999 (dark blue line) [Groeneveld, 2005] and 1000 (red line) [Groeneveld et al., 2008]. The δ¹⁸O records of the thermocline dweller N. dutertrei dextral from sites 999 (light blue line) and 1000 (orange line) and Mg/Ca temperature records of N. dutertrei (dextral variety) from sites 999 (thick dark blue line with dots) and 1000 (thick red line with dots).
The early Pliocene formation of the modern Caribbean Warm Pool more than compensated for the effect of AMOC-induced thermocline shoaling on Caribbean TCD.

4.5. Development of the EEP Cold Tongue

While the temperature evolution at the bottom of the photic zone in the tropical eastern Pacific was synchronous across the equator (Figure 4), a meridionally different de-
The development of surface hydrography between the doldrums and the EEP cold tongue may reflect regional differences in wind stress. The similar mixed layer temperature evolution (derived from Mg/Ca G. sacculifer) at doldrum Site 1241 [Groeneveld et al., 2006] and at Site 847 [Wara et al., 2005], today located at the northern edge of the EEP cold tongue region [Wara et al., 2005], suggests that both sites were located in warm surface water outside the EEP cold tongue region during the early to middle Pliocene (Figures 1 and 7b). Absolute $\delta^{18}$O G. sacculifer values and fluctuations are also similar at sites 1241, 847, and at ODP Site 851 (today located north of the modern EEP cold tongue region) [Cannariato and Ravelo, 1997; Wara et al., 2005; Steph et al., 2006b] (Figures 1 and 7c). Yet the pattern of long-term changes in $\delta^{18}$O records displays larger deviations from their associated Mg/Ca temperature records, especially during the earliest Pliocene (Figure 7). This suggests that the $\delta^{18}$O G. sacculifer signal at these sites is partly modulated by changes in mixed layer salinity.

Within the EEP cold tongue region, the development of a shallow thermocline between ~4.8 and 4.0 Ma should have increased the potential of trade winds to decrease SST by initiating coastal and equatorial upwelling. Indeed, the G. sacculifer $\delta^{18}$O record from equatorial upwelling Site 1239

![Figure 8.](image-url)

**Figure 8.** (a) Eastern equatorial Pacific meridional temperature gradient ($^\circ$C) estimated from the difference between the mean of the Mg/Ca-based SST reconstructions from sites 1241 and 847 (northeastern equatorial Pacific [Groeneveld et al., 2006; Wara et al., 2005]) and the alkenone-based SST from site 846 (southeastern equatorial Pacific [Lawrence et al., 2006]). All SST data were filtered with a three-point running mean, and the resulting time series were interpolated onto an equidistant grid with intervals of 25,000 years using a cubic spline. The strength of the meridional SST gradient determines the position of the ITCZ in the EEP, the strength of the cross-equatorial trade winds, and hence the efficiency of equatorial upwelling [Timmermann et al., 2007a]. (b) The comparison of $U_{37}^{K}$ temperatures from cold tongue Site 846 (blue) [Lawrence et al., 2006] with Mg/Ca G. sacculifer derived temperatures from West Pacific Warm Pool Site 806 (red) [Wara et al., 2005; Fedorov et al., 2006] indicates that the development of the equatorial Pacific east-west SST gradient started as early as ~4.0 Ma, with a second major step occurring at 1.7 Ma as previously suggested by, e.g., Fedorov et al. [2006].
diverges from the site 1241, 851, and 847 $\delta^{18}O_{G. sacculifer}$ records after ~3.9 Ma, and more clearly after ~3.7 Ma, displaying relatively higher $\delta^{18}O$ values (Figure 7c). This indicates a different development of mixed layer hydrography within the EEP cold tongue region, with lower temperatures (and/or higher salinity) compared to the East Pacific Warm Pool. A more faithful recorder of the SST development within the cold tongue is the $U^{HJ}_{37}$ SST record from Site 846 [Lawrence et al., 2006] (Figure 7d). Today, this site is located within the center of the EEP cold tongue, where the upwelling of cold waters is driven by Ekman divergence. The Site 846 SST estimates suggest that sea surface cooling within the EEP cold tongue region (~1°C/Ma) started between ~4.3 Ma and 3.6 Ma [Lawrence et al., 2006]. Moderate early sea surface cooling (beginning around 4 Ma) is also observed in the upwelling area off the California margin [e.g., Dekens et al., 2007; Liu et al., 2008; Brierley et al., 2009], but is neither reflected in the low-resolution $U^{HJ}_{37}$ SST record nor in the Mg/Ca$_{G. sacculifer}$ derived mixed layer temperature record from East Pacific Warm Pool Site 1241 (Figures 7b and 7d). Decreasing SST in the EEP cold tongue region accordingly led to strengthening of the meridional SST gradient between the northeastern (sites 847, 1241) and southeastern (Site 846) equatorial Pacific during the early Pliocene (Figure 8a). This finding is supported by our modeling results, which also indicate that intensification of the AMOC related to CAS closing resulted in lower SST in the southeastern equatorial Pacific, whereas SST in the northeastern equatorial Pacific increased (see Figure S1 in Text S1). The mechanisms involved in establishing this meridional SST gradient have been described in detail by Zhang and Delworth [2005], Timmermann et al. [2007b] and Xie et al. [2008]. They play a key role in establishing the annual cycle of SST on the equator and in modulating ENSO variability.

Comparing the $U^{HJ}_{37}$ SST record from EEP cold tongue Site 846 [Lawrence et al., 2006] with the Mg/Ca$_{G. sacculifer}$ temperature record from West Pacific Warm Pool Site 806 [Wara et al., 2005; Fedorov et al., 2006] furthermore suggests the initial development of an equatorial Pacific east-west SST gradient around 4.0 Ma (Figure 8b), consistent with early cooling observed in the EEP cold tongue region [e.g., Lawrence et al., 2006; Dekens et al., 2007]. In contrast, Fedorov et al. [2006] concluded that the transequatorial SST gradient was constantly weak during the time interval before ~1.7 Ma by comparing temperature records from Site 806 and tropical northeast Pacific Site 847. Yet the $U^{HJ}_{37}$ SST record from EEP cold tongue Site 846 might be better suited for assessing the Pliocene development of the transequatorial temperature gradient than Site 847, because the similarity of mixed layer temperatures at Site 847 and at East Pacific Warm Pool Site 1241 (Figure 7b) indicates that

Figure 9. Schematic relationships between AMOC-induced changes in TCD and wind stress with respect to the development of the modern EEP cold tongue (EEPCT). (a) Thermocline shoaling in the tropical eastern Pacific at 4.8–4.0 Ma yielded an important precondition for the turnaround from a warm eastern equatorial Pacific to the modern EEP cold tongue state. (b) Although EEP cold tongue sea surface cooling may have started as early as ~4.3–4.0 Ma ago [Lawrence et al., 2006] southeast trade winds were probably too weak to significantly amplify equatorial and coastal upwelling in a warm world prior to ~3.6 Ma. (c) Since 3.6–3.5 Ma, changes in the tropical wind field might then have tipped the scale for an increase in upwelling and productivity parallel to the intensification of NHG [Mudelsee and Raymo, 2005].
Site 847 was probably not located in the center of the EEP cold tongue region during the early and middle Pliocene.

[40] Changes in biogenic opal accumulation rates, a proxy for biological productivity, provide additional evidence for enhanced early Pliocene upwelling in the EEP cold tongue, since this region developed as the equatorial center of diatom productivity after ∼4.4 Ma [Farrell et al., 1995]. Diatom productivity at EEP cold tongue Site 1239 increased significantly around 3.6–3.5 Ma, when opal accumulation rates doubled (Figure 7e). This change coincides with an increase in the SST gradient between doldrum Site 1241 and EEP cold tongue Site 846 (Figure 7d) and may indicate both, enhanced upwelling or an increase in nutrient concentration, which might be related to a change in advective nutrient supply. We surmise that strengthening of the pole-to-equator temperature gradient in response to the intensification of NHG enhanced trade wind strength [Timmermann et al., 2004] and low-latitude upwelling since 3.6 Ma. If productivity changes are considered as additional evidence for the development of the EEP cold tongue, it occurred at least as early as 3.6–3.5 Ma.

5. Summary and Conclusions

[41] Both the early thermocline shoaling in the tropical eastern Pacific (4.8–4.0 Ma) and the early start of EEP cold tongue cooling (between ∼4.3 Ma and 3.6 Ma) [e.g., Lawrence et al., 2006; Dekens et al., 2007] suggest a chain of forcing mechanisms for the development of the modern EEP cold tongue, which is different from the well-cited hypothesis that this Pliocene climate switch was primarily linked to the major intensification of NHG (after ∼3.0 Ma) [e.g., Wara et al., 2005; Fedorov et al., 2006]. Instead, the timing of these events suggests a close link to the gradual closure of the CAS, leading to the development of the modern Atlantic–Pacific salinity contrast and to the amplification of AMOC [e.g., Keigwin, 1982; Haug and Tiedemann, 1998; Haug et al., 2001; Steph et al., 2006a].

[42] Although the impact of the gradual CAS closure on global climate change has recently been questioned [Molnar, 2008], our study brings the CAS back into play by providing for the first time a consistent link between the observed decrease in tropical east Pacific TCD and an increase in the strength of the AMOC during the early Pliocene (4.8–4.0 Ma) that was triggered by a threshold in the closure history of the CAS (Figure 9). Our new Caribbean proxy data also suggests that this threshold in the CAS closure led to the formation of the modern Caribbean Warm Pool (starting at ∼4.8 Ma), which more than compensated for AMOC-induced shoaling of the Caribbean thermocline. In contrast to previous studies, we identify changes in AMOC as a potential forcing mechanism, which consistently explains the Pliocene long-term evolution of tropical east Pacific TCD as well as the early onset of sea surface cooling in the EEP cold tongue [e.g., Lawrence et al., 2006; Dekens et al., 2007] and the development of a meridional SST gradient in the EEP during the early Pliocene (Figure 9).

[43] Further support for our main conclusions comes from a recent modeling study by Lunt et al. [2008]. Using the HadCM3 model, these authors find that closure of the CAS leads to strengthening of the AMOC as well as to warming of the Northern Hemisphere and cooling of the Southern Hemisphere (Figure 10a). This SST response to CAS closure is very similar to the SST response of the HadCM3 model to an intensification of the AMOC [Timmermann et al., 2007b] (Figure 10b) and to the ECBILT-CLIO SST response to CAS closing (see auxiliary material). The enhancement of the EEP cold tongue results from a northward shift of the ITCZ and an associated southeasterly trade wind intensification south of the equator, while the wind-evaporation-SST feedback helps to maintain this feature. Summarizing, independent modeling and paleo-proxy evidence suggests that the CAS played a key role in determining the strength of the AMOC, the position of the

Figure 10. (a) SST response (°C) to closing of the CAS (from Lunt et al. [2008] with kind permission of Springer Science and Business Media) as simulated by the HadCM3 coupled model. (b) SST response (°C) to intensification of the AMOC as simulated by the HadCM3 model [Timmermann et al., 2007b; Dong and Sutton, 2007].
ITCZ, the strength of the meridional SST gradient in the EEP and the existence of the EEP cold tongue.

The observed increase in biogenic productivity and upwelling within the EEP cold tongue after 3.6–3.5 Ma likely resulted from enhanced trade wind strength due to strengthening of the pole–equator temperature gradient in response to the intensification of NHG (Figure 9). Yet the thermocline deepening at doldrum Site 1241 after 3.0 Ma was presumably linked to AMOC weakening in response to the intensification of NHG. This contradicts the hypothesis that cooling of the deep ocean after 3.0 Ma led to a global decrease in tropical TCD.

A recent analysis of IPCC model results examining the next 100 years indicates a thermodynamically driven weakening of the Walker circulation in response to global warming, possibly favoring El Niño–like climate conditions [Vecchi and Soden, 2007]. Our results further suggest changes in AMOC as an important factor for the assessment of future changes in tropical climate dynamics. If global warming led to a reduction in both AMOC strength and trade wind intensity, the consequent increase in TCD would support the disappearance of the EEP cold tongue. Whether this mechanism may be adopted to assess future changes in ENSO variability remains to be shown.

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References
Keigwin, L. D. (1978), Pliocene closing of the Isthmus of Panama, based on ostracodographic


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