is less than 100%). Models for the additional oligonucleotide, GTP molecules and  $Mg^{2+}$  ions, have been fitted into electron density maps and refinement of these oligo— $Mn^{2+}$ —polymerase and oligo—GTP-Mg-Mn—polymerase complexes against their data sets, imposing strict threefold NCS constraints, resulted in models with R factors of 23.7 and 21.4%, respectively, and good stereochemistry (Table 1).

#### **Figures**

Unless otherwise stated figures were drawn using BOBSCRIPT $^{26}$  and rendered with RASTER3D $^{27}$ .

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Correspondence and requests for materials should be addressed to D.I.S. (e-mail: dave@strubi.ox.ac.uk). Coordinates have been deposited in the RCSB Protein database under accesssion codes: 1HHS, 1HHT, 1HI0, 1HI1, 1HI8.

# correction

# Improved estimates of global ocean circulation, heat transport and mixing from hydrographic data

Alexandra Ganachaud & Carl Wunsch

Nature 408, 453-457 (2000).

In this first paragraph of this paper, the uncertainty on the net deep-water production rates in the North Atlantic Ocean was given incorrectly. The correct value should have been  $(15 \pm 2) \times 10^6 \, \mathrm{m}^3 \, \mathrm{s}^{-1}$ .

## errata

# Changes in Greenland ice sheet elevation attributed primarily to snow accumulation variability

J. R. McConnell, R. J. Arthern, E. Mosley-Thompson, C. H. Davis, R. C. Bales, R. Thomas, J. F. Burkhard & J. D. Kyne

Nature 406, 877-879 (2000).

As the result of an editing error, the 1993–1998 aircraft-based altimetry surveys of the southern Greenland ice sheet reported by Krabill *et al.* (1999) were erroneously described as satellitebased.

# Genome sequence of enterohaemorrhagic *Escherichia coli* 0157:H7

Nicole T. Perna, Guy Plunkett III, Valerie Burland, Bob Mau, Jeremy D. Glasner, Debra J. Rose, George F. Mayhew, Peter S. Evans, Jason Gregor, Heather A. Kirkpatrick, György Pósfai, Jeremiah Hackett, Sara Klink, Adam Boutin, Ying Shao, Leslie Miller, Erik J. Grotbeck, N. Wayne Davis, Alex Lim, Eileen T. Dimalanta, Konstantinos D. Potamousis, Jennifer Apodaca, Thomas S. Anantharaman, Jieyi Lin, Galex Yen, David C. Schwartz, Rodney A. Welch & Frederick R. Blattner

Nature 409, 529-533 (2001).

The Genbank accession number for the annotated sequence given in this paper was typeset incorrectly. The correct accession number is AE005174.

of 3D periodic mesostructured materials without assuming any structural models. The resolution for the structure is primarily limited by the quality of the HREM images, which depends on the long-range mesoscale ordering. Therefore, although further progress may give better resolution, we expect no future change to the present conclusions about the structures of SBA-1, SBA-6 and SBA-16, because the validity of the solutions does not depend on the resolution. This is a characteristic of our method that makes it different from other approaches. We also suggest that the results presented here provide a quantitative topological description of ordered mesostructured composites, and that such descriptions are essential in understanding the properties and possible applications of the composites. The resolution of periodically ordered, 3D arrangements of bimodal (meso-micro) pores in SBA-1 and SBA-6 makes it possible to consider the detailed characterization of the range of complicated porous phases that are now synthetically achievable.

### Methods

# Synthesis of SBA-6

3.75 g of tetraethoxysilane (TEOS) was added with magnetic stirring to a clear solution containing 0.5 g of the gemini surfactant  $18B_{4\cdot3-1}$  (N,N,N,N'N'-pentamethyl-N'-  $[4\cdot(4\text{-octadecyloxyphenoxy})\text{-butyl}]\text{-propane-1,3-diammonium dibromide,} $C_{18}H_{37}\text{OC}_6H_4\text{OC}_4H_8\text{N}(\text{CH}_3)_2\text{C}_3H_6\text{N}(\text{CH}_3)_3\text{Br}_2),}$ 45.4 g of doubly distilled water, and 3.69 g of benzyltrimethylammonium hydroxide at room temperature. Stirring was continued for 20 h after the addition of TEOS at room temperature. The reaction gel mixture was heated for 2 d at 80 °C without stirring. The precipitate was filtered and dried in air at room temperature.$ 

#### **Determination of properties**

Ar adsorption and desorption isotherms were measured at 87 K. Pore volumes (cm $^3$  g $^{-1}$ ) for SBA-1, SBA-6 and SBA-16 are 0.6, 0.86 and 0.45, respectively, and the ratios of the pore volume to unit cell are respectively 0.57, 0.65 and 0.47. The surface-area/pore-volume ratio (2.26  $\times$  10 $^9$  m $^{-1}$ ) for SBA-1 is nearly three times that of SBA-6 (7.93  $\times$  10 $^8$  m $^{-1}$ ). The silica wall densities determined with an AccPyc 1300 helium pycnometer are also substantially different for SBA-1 (2.00 g cm $^{-3}$ ) and SBA-6 (2.20 g cm $^{-3}$ ).

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# Improved estimates of global ocean circulation, heat transport and mixing from hydrographic data

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Through its ability to transport large amounts of heat, fresh water and nutrients, the ocean is an essential regulator of climate<sup>1,2</sup>. The pathways and mechanisms of this transport and its stability are critical issues in understanding the present state of climate and the possibilities of future changes. Recently, global high-quality hydrographic data have been gathered in the World Ocean Circulation Experiment (WOCE), to obtain an accurate picture of the present circulation. Here we combine the new data from high-resolution trans-oceanic sections and current meters with climatological wind fields, biogeochemical balances and improved a priori error estimates in an inverse model, to improve estimates of the global circulation and heat fluxes. Our solution resolves globally vertical mixing across surfaces of equal density, with coefficients in the range  $(3-12) \times 10^{-4} \,\mathrm{m}^2 \,\mathrm{s}^{-1}$ . Net deepwater production rates amount to  $(15 \pm 12) \times 10^6 \,\mathrm{m}^3 \,\mathrm{s}^{-1}$  in the North Atlantic Ocean and  $(21 \pm 6) \times 10^6 \,\mathrm{m}^3 \,\mathrm{s}^{-1}$  in the Southern Ocean. Our estimates provide a new reference state for future climate studies with rigorous estimates of the uncertainties.

Obtaining a consistent picture of the oceanic circulation requires adjusting thousands of parameters consistently with a priori error estimates. We present here our best estimate from selected hydrographic data (Fig. 1), which will improve with the appearance of new data. Mass flux is the most basic element of the circulation and Fig. 2 shows the best-estimate coast-to-coast integrated water mass transports for selected density classes. A volume of  $15 \pm 2 \, \text{Sv}$  $(1 \text{ sverdrup} = 1 \times 10^6 \text{ m}^3 \text{ s}^{-1})$  of North Atlantic Deep Water (NADW) is produced in the northern North Atlantic Ocean and moves southward, entraining Antarctic Bottom Water (AABW) from below, and Antarctic Intermediate Water (AAIW) from above. As a result, the NADW is increased to  $23 \pm 3$  Sv as it exits the South Atlantic at 30°S. In the Southern Ocean, a total of  $21 \pm 6$  Sv of bottom water is formed from lower Circumpolar Deep Water (CDW)—which corresponds approximately to the lower NADW density range. Bottom water (NADW + AABW mixture) to the Atlantic, Indian and Pacific oceans are  $6 \pm 1.3$  Sv,  $11 \pm 4$  Sv and  $7 \pm 2$  Sv, respectively. In the Indian and Pacific oceans, most of this water returns southward at deep and intermediate levels. These net values are the sums of large, strongly spatially varying, flows of opposing sign, and thus oversimplify the actual circulation; a detailed description of the circulation within each ocean basin will be published elsewhere<sup>3,4</sup>. Our standard model estimate of the inflow in the South Pacific Ocean is in the lower range of previously published values, but it depends directly upon the weight given to the "PO" phosphate-oxygen combination (see Methods<sup>4,5</sup>) conservation constraints relative to mass conservation<sup>3</sup>. The deep inflow to the North Pacific Ocean is also weaker than previously found<sup>5</sup>, as a consequence of our consideration of heat and salt conservation in the northern parts of those basins.

No definition of bottom-water formation can be completely unambiguous because of the entrainment of ambient fluid during the sinking process. In our Southern Ocean definition, the bottom-

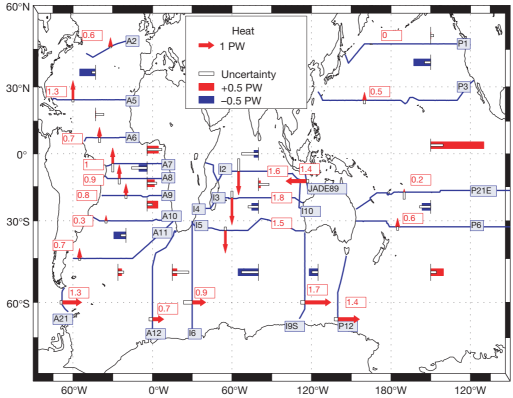
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water formation rate corresponds to the total amount of water crossing the neutral surface,  $\gamma^n = 28.11 \,\mathrm{kg} \,\mathrm{m}^{-3}$  downwards, and subsequently ventilating the deep Atlantic, Indian and Pacific basins. This water is provided primarily from the density range of lower NADW, so that the two sources of deep water are not additive, as is often assumed in qualitative calculations<sup>6</sup>. Consistent with conventional wisdom, no very large amount of deep water forms in the North Pacific or Indian oceans, although intermediate waters are produced there. Transient tracer measurements suggest AABW formation rates of water denser than  $\gamma^n = 28.27 \text{ kg m}^{-3}$  at 5 to 8 Sv (ref. 7). Because this density class is almost immediately mixed with CDW above it when exiting the Southern Ocean, it is not separately resolved in our model, and there is no inconsistency. However, there is a contradiction with the suggestion that the current production rate of AABW recently decreased<sup>6</sup>, as we find no inconsistency between the deep, contemporary, geostrophic circulation and the interior tracer field which reflects the circulation integrated over hundreds of years. We believe that the origin of the disagreement is in those authors' ad hoc assumptions<sup>6</sup> about the originating water masses.

The integrated circulation in the upper layers is more uncertain (Fig. 2, red arrows) owing to the enhanced temporal variability known to be present there<sup>3</sup>. A northward flow of  $16 \pm 3$  Sv of thermocline water from the South Atlantic Ocean balances the NADW southward flow; the strength of the Pacific-Indonesian throughflow is estimated at  $16 \pm 5$  Sv, consistent with the most recent current-meter measurements<sup>8</sup>. This estimate is sensitive to

the use of hydrography at very low latitudes where noise susceptibility and seasonal variability are large.

Vertical exchanges (advection, denoted  $w^*$ , and mixing, measured by a diffusion coefficient  $\kappa^*$  across the neutral surfaces<sup>9</sup> defining the layers) are associated with the general circulation. Both parameters are now the focus of intense research10,11 because numerical models are known to be very sensitive to  $\kappa^*$ . Direct measurements have shown little mixing in the ocean interior, while enhanced mixing of order 1,000 cm<sup>2</sup> s<sup>-1</sup> has been observed in oceanic regions with rough topography<sup>11</sup>. Table 1 summarizes the abyssal dianeutral (near vertical) exchanges by basin and depth range. Let an overbar denote a basin average. Then, in the deep range (=2,000-3,500 m), dianeutral velocities are  $\overline{w^*} = (0.1-0.6) \times 10^{-6}$  m s<sup>-1</sup>, and the diffusivities are  $\overline{\kappa^*} = (3-4) \times 10^{-4} \,\mathrm{m^2 \, s^{-1}}$  with global of  $\overline{w^*} = (0.13 \pm 0.03) \times 10^{-6} \,\mathrm{m \, s^{-1}}$  $(3.7 \pm 0.7) \times 10^{-4}$  m<sup>2</sup> s<sup>-1</sup> between 30° S and 47° N. In the bottom layers (=3,800 m to the bottom), both means are larger:  $\overline{w^*} = (0.4 \pm 0.1) \times 10^{-6} \,\mathrm{m \, s^{-1}}$  and  $\overline{\kappa^*} = (9 \pm 2) \times 10^{-4} \,\mathrm{m^2 \, s^{-1}}$ . Diffusivity values in the deep range are consistent with previous calculations from a crude one-dimensional global balance<sup>10</sup>. The largest values are found in the bottom layers although they are correspondingly more uncertain<sup>3</sup>. The spatial average values that are required by our tracer balance result from all mixing processes, including probable strong mixing generated near topography<sup>10,11</sup>. We interpret those values<sup>10</sup> as being the basin-average value of a process strongly mixing the ocean at highly localized regions.



**Figure 1** Hydrographic sections and heat fluxes. Transoceanic sections from the WOCE program were selected to ensure a reasonable temporal consistency (1990–1996) and to avoid crossing sections. Between Jakarta and Australia, the 1989 section from the Franco-Indonesian JADE program was used while pre-WOCE sections (1985, 1987)<sup>3</sup> were used in the North Pacific and at 32°S in the Indian Ocean. Each section is a collection of high-density temperature, salinity, oxygen and nutrient measurements. From temperature and salinity, a geostrophic velocity field is calculated and adjusted so that mass and other conservative tracers (see Methods) are conserved between sections. The resulting heat (or 'enthalpy', where the net mass flux is non-zero) transports are indicated

by the arrows and red numbers (positive northward/eastward). The white box at the tail end of each arrow is the one-standard-deviation uncertainty. Between sections, ocean—atmosphere heat transfers are indicated by the zonal length of the coloured boxes (blue for ocean cooling; red for ocean heating), with the length of the white box inside indicating the uncertainty. (Because the ocean—atmosphere heat transfers are anomaly residuals, that is, corrected for residual mass imbalances, they do not correspond exactly to the differences between net fluxes across sections, for example, in the North Indian Ocean. But this discrepancy is much less than the uncertainties.)

Diffusivities could not be resolved in the Southern Ocean, where many neutral surfaces outcrop. The improved inverse model method has produced the first near-global, resolved estimates of the dianeutral transfers. The overall results are inconsistent with recent suggestions that the ocean mixes primarily at near-surface outcrops of the neutral surfaces, that is, primarily in the Southern Ocean<sup>12</sup>. Strong abyssal mixing is required by the observed geostrophically balanced circulation, and its absence is incompatible with the observed property distributions.

Figure 1 shows the heat (actually, enthalpy) transports, across each hydrographic section (arrows) along with the residuals reflecting atmospheric heat exchanges (boxes). Residuals are accurately determined at middle and high latitudes, but are more uncertain at lower latitudes (for example, in the Atlantic Ocean) owing to an enhancement of the geostrophic noise there<sup>3</sup>. Nevertheless, the total heating over the tropical Atlantic and Pacific oceans are welldetermined, respectively  $0.7 \pm 0.2 \,\mathrm{PW}$   $(1 \,\mathrm{PW} = 10^{15} \,\mathrm{W})$  and  $1.6 \pm 0.4 \,\mathrm{PW}$ . No significant heat transfers are found in the Indian Ocean because of the large, uncertain, warm water inflow from the Pacific Ocean. This large warm water flux is the main heat escape from the Pacific Ocean, resulting in a northward heat flux in the South Pacific. In the southern Pacific sector, significant heating is found, in contrast with the sparse in situ observations<sup>13</sup>, but in qualitative agreement with the recent re-analysis of the European Centre for Medium Range Weather Forecasts<sup>14</sup>. Figure 3 shows the globally integrated heat fluxes compared to independent estimates. Most of the cooling occurs in the Northern Hemisphere, at a rate of  $-1.7 \pm 0.2$  PW, in balance with the  $2.3 \pm 0.4$  PW heating in the

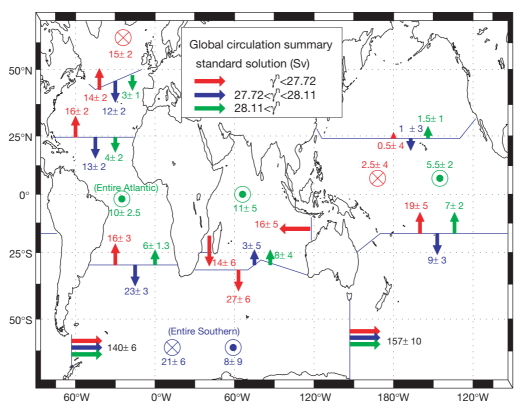
tropical band and the  $-0.7 \pm 0.3 \, \mathrm{PW}$  cooling in the Southern Ocean

Changes in the oceanic heat transport can have a large impact on atmospheric temperature gradients<sup>15,16</sup> and thus on climate. Previous estimates of the ocean–atmosphere heat exchanges that are based upon purely ocean surface observations are highly uncertain<sup>17,18</sup>. Analyses from numerical weather prediction centres provide oceanic surface fluxes that are often used as boundary conditions for driving ocean models, but associated uncertainty estimates are not provided. Heuristic calculations suggest uncertainties in their estimates of at least  $\pm 0.6 \, \mathrm{PW}$  for the meridional oceanic heat transport at most latitudes<sup>19</sup>. The present inversion indicates uncertainties that depend on latitude, with a high accuracy of globally integrated heat transfers (Fig. 3). Similar budgets, to be

Table 1 Basin-averaged dianeutral velocities and diffusivities

	$\overline{w} * (10^{-6} \mathrm{m  s^{-1}})$	$\overline{\kappa} * (10^{-4} \mathrm{m}^2 \mathrm{s}^{-1})$
Atlantic bottom	0.5 ± 0.2	9 ± 4
Indian bottom	$0.6 \pm 0.3$	$12 \pm 7$
Pacific bottom	$0.4 \pm 0.1$	9 ± 2
Southern bottom	$-0.25 \pm 0.1$	_
Atlantic deep	$0.1 \pm 0.05$	3 ± 1.5
Indian deep	$0.3 \pm 0.15$	4 ± 2
Pacific deep	$0.1 \pm 0.03$	4 ± 1
Southern deep	$0.1 \pm 0.1$	_

The average is calculated on neutral surfaces from  $\gamma^0=28.1$  kg m $^{-3}$  to the bottom (generally 3,800 decibars (or metres) to the bottom) for the 'bottom layers' and from  $\gamma^0=27.96$  to  $\gamma^0=28.07$  for the 'deep layers' (generally 2,000 m to 3,500 m).



**Figure 2** Zonally integrated layer mass transports. The estimated water transports are indicated for the different density classes bounded by neutral surfaces ( $\gamma^n$ , in kg m<sup>-3</sup>) and across selected hydrographic sections. Neutral surfaces are close to surfaces of constant density, but are chosen so that movement along them minimizes the work done against gravity. The colour of the upwelling or down-welling arrows indicates the layer from which the water is coming. A flux of 0.8 Sv from the Pacific to the Atlantic Ocean through the Bering Strait was taken into account, although it is much smaller than the uncertainties on

the net transports. In the Southern Ocean, the bottom water formation takes place mostly in the Weddell Sea, while the upwelling distribution is uncertain. In the Indian Ocean, most of the upwelling takes place north of  $7^{\circ}$  S. The South Pacific transports are given at  $17^{\circ}$  S because of the more complicated structure at  $32^{\circ}$  S (ref. 3). Note the increase in the Southern Ocean transport south of Australia owing to the recirculation of Indonesian throughflow water.

published elsewhere, of oxygen and nutrients, are important for both climate and biogeochemical processes<sup>3,20</sup>.

The physical circulation estimated here during WOCE provides a new reference state for climate change studies, and a test-within the estimated uncertainties—of general circulation models of both atmosphere and ocean. The estimated values of  $\kappa^*$  and  $w^*$  represent values that models will need to simulate. The high-resolution WOCE data set, the confinement in time, the use of currentmeter data as constraints in boundary currents, and the detailed analysis of the uncertainties associated with our method are great improvements in reliability over previous circulation estimates. Within the error estimates, there is no conflict in either transports or heat fluxes with a previous global inversion that was based on a nearly independent and coarser data set<sup>1,21</sup>, and no statistically significant change in integrated mass transports over the past 30 years was found<sup>1,22</sup>. This is an important conclusion in its own

Improving the accuracy of the estimates made from the present data sets, if interpreted as climatological averages, will not be easy. Although the addition of new data, for example, existing or future meridional sections, will better the spatial resolution, limitations now lie primarily in the uncertainty introduced by true oceanic variability from the daily to the interannual. Significant improvements in the present numbers will occur only through the use of data sets permitting true temporal averaging of the oceanic circulation. For instance, the present solution has large uncertainties due to undersampling of the highly variable Brazil current and Pacific-Indian throughflow. Additional observations there would greatly improve accuracy. Further refinement of the a priori error estimates

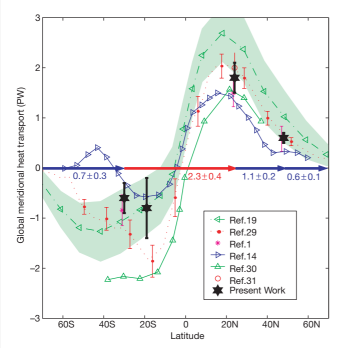


Figure 3 Global meridional heat transports. The black stars with thick error bars indicate the global, zonally integrated heat transports. Ocean-atmosphere heat exchanges between selected latitudes are given by numbers above the northward pointing arrows (positive for oceanic heating). The uncertainties from ref. 19 are indicated by the green band. The previous estimate of ref. 1 is different, but statistically consistent. No incompatibility is found between the estimates by the European Centre for Medium Range Weather Forecasts and US National Center for Environmental Prediction models 14,19,29 and our new values, owing to their large uncertainties<sup>19</sup>. Other climatological estimates<sup>30,31</sup> were published without error estimates, thus precluding any quantitative comparison. The estimate of ref. 30 is taken from their experiment B. (This figure is adapted and completed from refs 1 and 30.)

by using improved numerical models and other data sets (such as altimetry) is also possible.

# Methods

The method is that of hydrographic inverse box models<sup>23</sup>, where the relative, geostrophic velocity field is obtained from temperature and salinity measurements across the sections of Fig. 1. This initial flow is uniformly adjusted at each location so that the flow satisfies near-conservation of mass, anomalies of salt, heat, and the phosphate-oxygen combination ("PO" =  $170[PO_4] + [O_2]$ )<sup>24,25</sup> within oceanic layers defined by neutral surfaces (neutral surfaces are densities chosen so that work against gravity is minimized when following the surface, permitting global use of a single density variable<sup>26</sup>). Silica conservation is also required, top-to-bottom, while heat and PO are conserved only in layers that are not in contact with the surface. The surface layer is directly wind-driven (Ekman transport). Diffusive and advective exchanges between layers are determined by the model. The use of anomaly equations<sup>27</sup> permits, through noise subtraction, determination of diffusivities across each interface. Substantial improvements were made to the method, compared to that used in ref. 1, with, in particular, rigorous estimates of uncertainties through a determination of the a priori model error based on a simulation from the output of a quasi eddy-resolving (1/4°) ocean general circulation

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# The use of earthquake rate changes as a stress meter at Kilauea volcano

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Stress changes in the Earth's crust are generally estimated from model calculations that use near-surface deformation as an observational constraint. But the widespread correlation of changes of earthquake activity with stress<sup>1-5</sup> has led to suggestions that stress changes might be calculated from earthquake occurrence rates obtained from seismicity catalogues. Although this possibility has considerable appeal, because seismicity data are routinely collected and have good spatial and temporal resolution, the method has not yet proven successful, owing to the nonlinearity of earthquake rate changes with respect to both stress and time. Here, however, we present two methods for inverting earthquake rate data to infer stress changes, using a formulation for the stress- and time-dependence of earthquake rates<sup>6</sup>. Application of these methods at Kilauea volcano, in Hawaii, yields good agreement with independent estimates, indicating that earthquake rates can provide a practical remote-sensing stress meter.

The inversions use a formulation for earthquake rate changes<sup>6</sup> derived from laboratory observations of rate- and state-dependent fault strength<sup>6–8</sup>, which constrain the earthquake nucleation process to be dependent on both time and stress. Previously, this formulation has been applied to model the spatial and temporal characteristics of earthquake clustering phenomena, including foreshocks and aftershocks<sup>6,7</sup>, and to evaluate earthquake probabilities following large earthquakes<sup>9</sup>. The effectiveness of the formulation for forward modelling of earthquake phenomena, and its derivation from observed fault properties, provide the basis for its use to estimate stress changes from earthquake rate data. This approach yields stresses that drive the earthquake process. As such, it is distinct from other seismological methods that yield measures of stress changes resulting from earthquakes.

The formulation of Dieterich<sup>6</sup> for rate of earthquake activity R (in a specified magnitude range) can be written in the condensed form

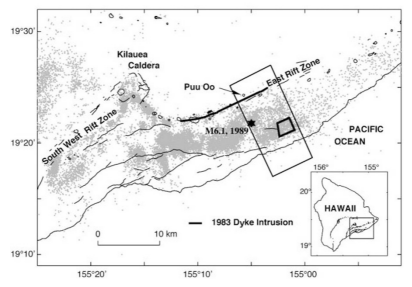
$$R = \frac{r}{\gamma \dot{S}_{r}}, \text{ where } d\gamma = \frac{1}{A\sigma} [dt - \gamma dS]$$
 (1)

where  $\gamma$  is a state variable, t is time, and S is a modified Coulomb stress function defined below. The constant t is the steady-state earthquake rate at the reference stressing rate  $\dot{S}_r$ . A is a dimensionless fault constitutive parameter with values usually in the range 0.005–0.015 (refs 6–8). The modified Coulomb stress function is defined as

$$S = \tau - [\mu - \alpha]\sigma \tag{2}$$

where  $\tau$  is the shear stress acting across fault planes that generate earthquakes (positive in the slip direction),  $\sigma$  is the normal stress (less pore fluid pressure),  $\mu$  is the coefficient of fault friction and  $\alpha$  is a constitutive parameter<sup>6,10</sup> with an assigned value in this study of 0.25 (refs 6, 10). In equation (1), the term  $A\sigma$  is a constant (that is, changes in  $\sigma$  are negligible relative to total  $\sigma$ ). For a stress step, equation (1) yields the characteristic aftershock sequence, which consists of an immediate jump of seismicity rate followed by decay that obeys the Omori  $t^{-1}$  aftershock decay law with aftershock duration  $t_{\sigma} = A\sigma/\dot{S}$  (ref. 6).

We use two methods to estimate stress changes from earthquake rate data. The first gives stress as a function of time in a specified volume. From equation (1), the observed rate R is used to directly calculate  $\gamma$  as a function of time (that is,  $\gamma(t) = r/R(t)\dot{S}_r$ ). This requires an estimate of  $\dot{S}_r$ , which can be obtained from independent



**Figure 1** Map of Kilauea volcano showing earthquakes of magnitude  $M \approx 1.5$  from 1976 to 1983. Intrusion of magma into the southwest and east rift zones of Kilauea causes rift expansion, and results in motion of the south flank to the SSE. Eruptive fissures initiated

the Puu Oo eruption, which started 1 January 1983 and continues to the present. The small polygon is the region of analysis of Fig. 2; the large rectangle gives the region of analysis of Figs 3 and 4.

# **Changes in Greenland ice sheet** elevation attributed primarily to snow accumulation variability

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The response of grounded ice sheets to a changing climate critically influences possible future changes in sea level. Recent satellite surveys over southern Greenland show little overall elevation change at higher elevations, but large spatial variability<sup>1-3</sup>. Using satellite studies alone, it is not possible to determine the geophysical processes responsible for the observed elevation changes and to decide if recent rates of change exceed the natural variability. Here we derive changes in ice-sheet elevation in southern Greenland, for the years 1978-88, using a physically based model of firn densification<sup>4</sup> and records of annual snow accumulation reconstructed from 12 ice cores at high elevation. Our patterns of accumulation-driven elevation change agree closely with contemporaneous satellite measurements of icesheet elevation change, and we therefore attribute the changes observed in 1978-88 to variability in snow accumulation. Similar analyses of longer ice-core records show that in this decade the Greenland ice sheet exhibited typical variability at high elevations, well within the long-term natural variability. Our results indicate that a better understanding of ice-sheet mass changes will require long-term measurements of both surface elevation and snow accumulation.

The previously reported, 1978-88 elevation-change estimates for the southern Greenland ice sheet1, derived from analysis of Seasat (6 July to 10 October 1978) and Geosat-Exact Repeat Mission (8 November 1986 to 7 November 1988) satellite radar altimeter data, were improved recently by adding data from the Geosat-Geodetic Mission (1 April 1985 to 30 September 1986). The updated analysis<sup>5</sup> included twice as many elevation-change measurements and expanded the spatial coverage by 50%. In addition, a precise global-ocean reference network created from four years of Topex/ Poseidon altimeter data was used to obtain improved corrections for altimeter radial orbit errors and measurement system biases. The updated elevation-change map includes about 90% of the area above the 2,000-m elevation contour south of 72.1°N (maximum latitude of Seasat/Geosat) and about 75% of the area west of the elevation divide between 1,700-2,000 m (Fig. 1). Effectively no coverage is available below 1,700 m because of the poor quality and limited quantity of the radar altimeter data near the ice-sheet edge<sup>5</sup>. The rate of elevation change (dH/dt) varies spatially from -24to +24 cm yr<sup>-1</sup> over distances as small as 200 km (note the strong dH/dt gradient around 66°N and the abrupt transition from positive to negative change at the elevation divide).

Between 1997 and 1999, ice cores that penetrated at least to a depth corresponding to 1978 were collected at ten locations around the southern ice sheet. All cores were analysed for a number of seasonally varying chemical species (hydrogen peroxide, calcium, ammonium, nitrate ions) using a continuous ice-core melter analysis system<sup>6</sup>. Two additional seasonally varying constituents, insoluble dust concentration and stable water-isotope ratios, were measured using discrete samples. In combination with field measurements of snow density, these multi-parameter glacio-chemical records were used to determine records of net annual snow accumulation at each site. Because of spatial variability in snow accumulation<sup>7</sup>, multiple cores were collected at four locations to identify regional-scale versus local-scale accumulation components<sup>8,9</sup>. Discrete measurements of hydrogen peroxide from a core drilled in 1988 were used to develop a record of annual accumulation at Dye 3 (65.2°N, 43.9°W)<sup>10</sup> and a 1778–1989 accumulation record was developed at Crete (71.1° N, 37.3° W) by combining published accumulation records<sup>11,12</sup>.

The density of near-surface firn layers increases with depth and, in response to changing snow accumulation rates and temperature, this increase with depth also varies with time. To derive changing ice-sheet elevation from measured time series of snow accumulation, we used a model that predicts the evolution of the densitydepth profile by solving coupled heat- and mass-transfer equations using a one-dimensional, lagrangian scheme, referenced to pressure coordinates4. The constitutive relationship for snow was calculated using rate equations for the various sintering mechanisms important in snow compaction<sup>13,14</sup>. The initial depth profiles of temperature and density were obtained from steady-state calculations at the mean temperature, mean accumulation rate and mean surface density. Boundary conditions include the temperature of the ice at depth, and surface histories of temperature<sup>15</sup>, accumulation

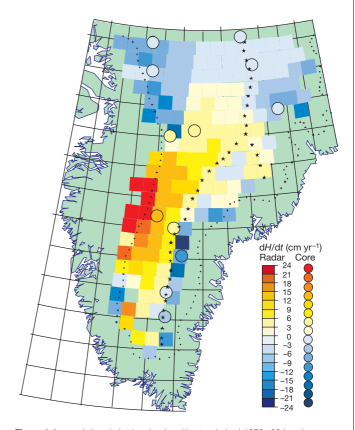


Figure 1 Accumulation-derived and radar-altimetry-derived 1978-88 ice-sheet elevation change. Shown are the approximate locations of the elevation divide (stars), the 2,000-m elevation contour (dots) and the ice-core sites (circles). The radar  $\mathrm{d}H/\mathrm{d}t$  values are shown in  $50 \times 50$  km cells (adapted from ref. 5).

rate and density. Mean surface density was derived from the ice-core data using the zero-depth intercept of a polynomial fit to the measured density—depth profile. Densification rates for the upper two metres were also derived from the measured density profiles rather than from sintering rate equations (at shallow depths the effects of vapour transport, seasonal temperature variations and reworking by wind are important although these were not explicitly included in the sintering model). The annual temperature cycle and surface density (uppermost 1 m) in Greenland change little from year to year. Earlier calculations<sup>4</sup> show that elevation changes resulting from these intra-annual variations are small compared with those resulting from inter-annual variations in snow accumulation.

The modelled change in vertical separation between the surface and the 4-MPa pressure level overburden (deep enough to lie within incompressible ice) defines the elevation change caused by variations in density<sup>4</sup>. The actual elevation change observed at any location is the sum of this component and the change in the local ice-equivalent-thickness (defined as the total mass of ice occupying a unit area of bedrock, divided by the density of pore-free ice). Because the divergence of the ice flow field at each core site is unknown, we cannot directly calculate the change in ice-equivalent thickness over the time period spanned by the ice core records. However, on the assumption that the ice-flow divergence is constant during this period, an estimate for the higher-frequency changes in ice-equivalent thickness can be obtained using the measured accumulation rates and added to the elevation changes predicted by the density model. We refer to the resulting time series as the 'elevation anomaly' at each site. These will be superimposed upon any secular trend in elevation change caused by long-term mass imbalance. Both the observed and accumulation-derived elevation changes at the twelve locations where annual accumulation data are available over the 10-year period are listed in Table 1 and shown in Fig. 1. To facilitate comparison with the radar-derived observations, 1978-88 accumulation-driven elevation change (dH/dt) is expressed in cm  $yr^{-1}$ .

We investigated the sensitivity in the calculation of accumulation-derived estimates of ice-sheet elevation change to several sources of error. To examine the sensitivity to errors in the constitutive equation for snow, strain rates in the densification model were artificially increased and decreased by a factor of ten<sup>4</sup>. To determine the sensitivity of our results to fluctuations in accumulation rate before the beginning of the ice-core records, we used an ensemble of six plausible ~50-year accumulation-rate histories having the same mean and inter-annual variance as the observed record in Monte-Carlo simulations. Similarly, we investigated the effects of short-scale (tens of metres) spatial variations of accumulation rate by adding noise to the measured time series and repeating these calculations for an ensemble of such records. The variance of the local noise component was obtained from closely

Table 1 Ice-core accumulation-derived and radar-based estimates of mean dH/dt from 1978 to 1988

Site latitude (° N), longitude (° W)	Initial year in accumulation record	1978 — 1988 Accumulation dH/dt (cm yr <sup>-1</sup> )	1978 — 1988 Radar* dH/dt (cm yr <sup>-1</sup> )
72.3, 38.0 (Summit)	1950	-1.1 (±2.9)	-0.1 (±1.5)
71.1, 37.3 (Crete)	1778	$-0.7 (\pm 4.2)$	$-1.9(\pm 1.5)$
71.9, 47.5	1974	-1.9 (±2.1)	$-8.0 (\pm 1.5)$
71.1, 47.2	1974	-1.1 (±2.0)	-5.1 (±2.1)
69.8, 35.0	1976	$-3.8 (\pm 1.5)$	$-6.2 (\pm 1.5)$
69.2, 43.0	1977	$+5.8 (\pm 1.6)$	$+4.5 (\pm 1.7)$
69.0, 45.0	1977	$+6.9 (\pm 2.3)$	$-22.3 (\pm 1.7)$
66.4, 46.2 (Dye 2)	1900	+12.7 (±2.3)	$+14.9 (\pm 1.6)$
66.0, 44.5	1976	$+4.8 (\pm 1.2)$	$+4.7 (\pm 1.6)$
65.2, 43.9 (Dye 3)	1931	$-10.3 (\pm 2.3)$	$-8.6 (\pm 1.8)$
63.8, 45.0	1978	$-0.4 (\pm 2.0)$	+3.7 (±1.6)
63.1, 44.8	1978	-3.2 (±2.0)	$-2.3 (\pm 1.8)$

Error limits show the 65% confidence interval.

\* Includes subtraction of 0.5 cm yr<sup>-1</sup> uplift from long-term isostatic rebound.

spaced ice cores at four of the ice-core sites<sup>8,9</sup>. Note that kilometrescale topography may also influence accumulation, although generally at multi-decadal to century, rather than annual, timescales. The 65% confidence intervals in Table 1 indicate the combined contribution from all these sources of error. Sensitivity to the accumulation history before the start of the core decreases rapidly with time under the relatively warm conditions found in Greenland<sup>4</sup>. Many of the core records extended to 1974 and earlier (Table 1) and for these sites, short-scale spatial variability in accumulation during the 1978-88 period provided the largest contribution to the combined uncertainties. At the six sites with shorter accumulation records, approximately half of the total uncertainty was the result of unknown accumulation before the start of the core record. If the previous accumulation were markedly different than during the core-based observation period, this would affect the results at sites with relatively short accumulation records.

The radar dH/dt values reported in Table 1 are the average of all elevation-change measurements within a circular area centred at the ice-core location. A nominal radius of 20 km was used for ten of the core locations, whereas a radius of 30 km was used at two core locations (Summit and Dye 3) where insufficient data were available within the nominal radius. Each set of data was edited by excluding elevation differences greater than two standard deviations from the primary gaussian distribution. The 65% confidence intervals include the combined contribution of random measurement error and estimates of the uncertainties in various corrections applied to the elevation data (isostatic uplift, inter-satellite bias, orbit-error corrections, and environmental corrections)<sup>1,5</sup>.

Comparisons between the accumulation-derived and radaraltimeter elevation-change estimates (see Supplementary Information) indicate that the large majority of the measured change in ice-sheet elevation in southern Greenland from 1978 to 1988 is the result of temporal variability in snow accumulation. The accumulation-derived elevation change gives the same general pattern of strong thickening to the west of the elevation divide and thinning to the east. We note the excellent match in the strong radar dH/dtgradient across the ice divide from Dye 2 (66.4° N, 46.2° W) to Dye 3 (65.2° N, 43.9° W). The notable exception is in the western region along the 2,000-m contour between 69°N and 72°N where the accumulation-derived elevation change is generally smaller than the radar-based observations. A significant difference was also found in this area in a comparison between 1978-88 radar dH/dt and 1993-98 laser dH/dt results<sup>16</sup> with no corresponding change in accumulation rate between the two time periods. Previous modelling and analysis of the Seasat and Geosat altimeter waveforms demonstrated that there was a significant temporal change in the geophysical characteristics of the near-surface snow in this particular area of the ice sheet<sup>17</sup>. Indeed, an unusually large number of melt events were found in the 69.0° N, 45.0° W core at depths corresponding to 1977 and 1978, with relatively few melt events in the early to mid-1980s. No such anomalous melt features were observed in the nearby 69.2° N, 43.0° W core where there is good

Table 2 Ice core accumulation-derived estimates of the standard deviation in decadal ice-sheet elevation change (dH/dt) over the specified time ranges

Latitude (°N), longitude (°W)	Time range	Standard deviation in decadal-scale dH/dt (cm yr <sup>-1</sup> )
78.5, 56.8	1690–1994	2.4
73.8, 49.5	1690-1994	4.1
72.3, 38.0	1950-1998	2.0
71.1, 37.3	1778-1982	3.7
70.3, 45.0	1177-1966	8.4
66.4, 46.2	1900-1998	12.7
65.2, 43.9	1931-1987	9.4
	78.5, 56.8 73.8, 49.5 72.3, 38.0 71.1, 37.3 70.3, 45.0 66.4, 46.2	78.5, 56.8 1690–1994 73.8, 49.5 1690–1994 72.3, 38.0 1950–1998 71.1, 37.3 1778–1982 70.3, 45.0 1177–1966 66.4, 46.2 1900–1998

1978–88 dH/dt values for Summit, Crete, Dye 2, and Dye 3 are given in Table 1.

agreement between the radar and accumulation dH/dt estimates. These observations indicate that a large melt layer existed in 1977 and 1978 which resulted in a different radar signature at the snowair interface than in subsequent years. This caused Geosat-Seasat dH/dt estimates to be biased towards excess thinning and/or reduced thickening in that region over the 10-year period.

To put these accumulation-derived changes in ice-sheet elevation in historical perspective, we derived estimates of dH/dt over longer time periods using previously published<sup>6,10,12</sup> and new records of annual accumulation. From the annual dH/dt time series, decadal dH/dt (cm yr<sup>-1</sup>) values were computed. Standard deviations in decadal dH/dt (Table 2) range from 2.0 cm yr<sup>-1</sup>at the Summit location in central Greenland to 12.7 cm yr<sup>-1</sup> at the southern Dye 2 site. Comparisons of the historical range of decadal dH/dt (Table 2) with the 1978-88 accumulation-derived and radarbased measurements (Table 1) indicate that recently observed changes in ice-sheet elevation above  $\sim 2,000$ -m elevation are typical. Because snow accumulation at Dye 2 is highly cyclic, even the marked increase in annual accumulation from 1978 to 1988 that has led to the recent large increase in ice-sheet elevation in that region is not unusual over the last century.

These widely distributed accumulation measurements and computed elevation anomalies demonstrate that altimetry-derived estimates of ice-sheet thickening and thinning from 1978-88 over much of the southern Greenland ice sheet above ~2,000-m elevation are consistent with elevation changes caused by temporal variability in snow accumulation. The ice-core accumulation measurements reported here do not address the significance of observed elevation change at lower elevations<sup>2</sup>. Analyses of longer accumulation records indicate that the decadal-scale changes in ice-sheet elevation that occurred during 1978-88 are typical over the last few centuries and well within the natural variability of accumulationdriven elevation change. It is clear that decadal-scale variability in snow accumulation has the potential to mask longer-period changes in surface elevation associated with ice-sheet dynamics. This suggests that accurate detection of any long-term mass imbalance of the ice sheets and assessment of likely causes will require multi-decadal time series of surface elevation in conjunction with widely distributed ice-core-derived accumulation measurements collected over the time period of interest.

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# **Evidence for iron, copper and zinc** complexation as multinuclear sulphide clusters in oxic rivers

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The availability and toxicity of trace metals in fresh water are known to be regulated by the complexation of free metal ions with dissolved organic matter<sup>1-3</sup>. The potential role of inorganic sulphides in binding trace metals has been largely ignored because of the reduced persistence of sulphides in these oxic waters. However, nanomolar concentrations of copper and zinc sulphides have been observed in four rivers in Connecticut and Maryland<sup>4,5</sup>. Here we report dissolved (< 0.2 µm particle diameter) sulphide concentrations ranging up to 600 nM, with more than 90% being complexed by copper, iron and zinc. These complexes account for up to 20% of the total dissolved Fe and Zn and 45% of the total dissolved Cu. Fourier transform mass spectrometry reveals that these complexes are not simple M(HS)<sup>+</sup> protonated species<sup>6,7</sup> but are higher-order unprotonated clusters (M<sub>3</sub>S<sub>3</sub>, M<sub>4</sub>S<sub>6</sub>, M<sub>2</sub>S<sub>4</sub>), similar to those found in laboratory solutions<sup>8-10</sup> and bio-inorganic molecules11. These extended structures have high stability constants<sup>8,10</sup> and are resistant to oxidation and dissociation<sup>10,12</sup>, which may help control the toxicity of these and other less abundant, but more toxic, trace metals, such as silver, cadmium and mercury.

We found the total dissolved sulphide (DS<sub>T</sub>) concentrations and metal sulphide speciation in river waters to be strongly correlated with the extent of watershed development. DS<sub>T</sub>, defined as  $\Sigma$  Fe sulphides + Cu sulphides + Zn sulphides + other soft class B metal sulphides (Ag, Cd, Hg, Pb) +  $HS^- + S_x^{2-} + S^0$ , was found in concentrations ranging between 20 and 580 nM (Table 1). High DS<sub>T</sub> (> 200 nM) was found in rivers that drained urbanized areas that received inputs of treated sewage effluent. In these rivers, the metal sulphide speciation, as determined by electrochemical analysis<sup>4,5,13,14</sup>, consisted of metal sulphide complexes in the following order of concentration abundance: Fe sulphides  $(\Sigma \text{ FeS} + \text{FeSH}^+) \gg \text{Zn sulphides} > \text{Cu sulphides} > (\text{polysulphides}).$ Polysulphides  $(S_{4,5}^{2-})$ , which are partly oxidized sulphide species that are reduced by Cr(II), were only electrochemically measured<sup>15</sup> in low examine the quaternary complex, LexA-AP3 and PI were expressed on the bait vector, and GAL4 AD-AG and/or SEP3-MIK were expressed on the prey vector. When two genes were expressed on the same vector, they were both driven by ADH1 promoters. Amino-acid residues 1-167 and 1-171 were used for the truncated AP1-MIK and SEP3-MIK proteins, respectively. Other processes and the colony-lift  $\beta$ -gal assays were performed in accordance with the manufacturer's instructions (Clontech).

#### **Immunoprecipitation**

For immunoprecipitation experiments, radiolabelled AP1 or SEP3 were mixed with haemagglutinin (HA)-tagged proteins and precipitated with anti-HA antibody. Precipitated AP1 and SEP3 were separated by SDS-PAGE and detected by radio-imaging analyser, BAS2000 (Fujifilm). Other procedures were done as described<sup>7,12</sup>.

#### Transactivation assay

For yeast, MADS proteins cDNAs were fused in-frame to GAL4 DNA-binding domain on pAS2-1 (Clontech) and transformed into the yeast strain YRG-2 (UAS::lacZ, Stratagene). AP1-K2C (residues 125-256) and SEP3-K2C (128-257) were used as truncated MADS proteins. Yeast cells were grown at 22 °C overnight, and the β-gal activity was assayed at 30 °C using o-nitrophenyl-β-D-galactopyranoside.

For onion epidermal cells, 35S promoter-driven MADS cDNAs that express native MADS proteins (effector) and CArG::LUC (reporter) were co-transfected into onion epidermal cells by using a particle delivery system (Bio-Rad). CArG::LUC has seven repeats of MADS protein binding consensus sequence<sup>29</sup>, 5'-GGGGTGGCTTTCCTTTTTGG TAAATTTTGGATCC-3' (CArG box is underlined), upstream of the 35S minimal promoter (-30). 35S::Renilla luciferase (RLUC) was used for the internal control. LUC assays were conducted using Dual-luciferase reporter system (Promega). Other procedures were done as described<sup>30</sup>.

#### Plant material

Arabidopsis Columbia ecotype was used for Agrobacterium-mediated vacuum transformation31. Plant crossing was carried out by manual cross-pollination. The presence of the transgenes was confirmed by PCR. AP3::GUS plants have a 600-base-pair region of the AP3 promoter<sup>16</sup>. Staining for GUS activity was done as described<sup>16</sup>.

#### Cryo-scanning electron micrograph

We used a Hitachi S-3500N scanning electron microscope equipped with a cryo-stage. For observation and photography, the stage was chilled at −20 °C and the natural scanning electron microscopy (SEM) mode (70 Pa) was used with a 25-kV accelerating voltage.

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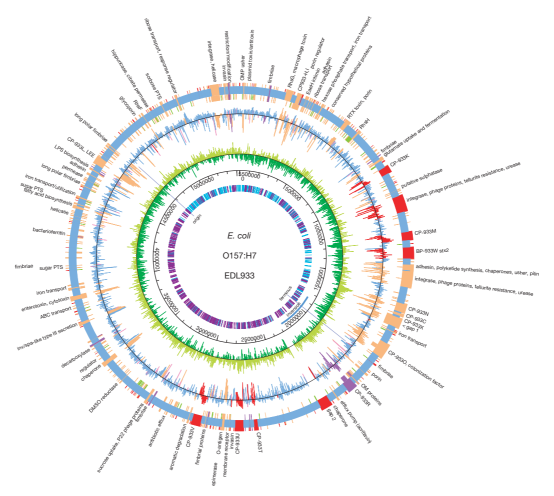
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# **Genome sequence of** enterohaemorrhagic Escherichia coli 0157:H7

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The bacterium Escherichia coli O157:H7 is a worldwide threat to public health and has been implicated in many outbreaks of haemorrhagic colitis, some of which included fatalities caused by haemolytic uraemic syndrome<sup>1,2</sup>. Close to 75,000 cases of O157:H7 infection are now estimated to occur annually in the United States<sup>3</sup>. The severity of disease, the lack of effective treatment and the potential for large-scale outbreaks from contaminated food supplies have propelled intensive research on the pathogenesis and detection of E. coli O157:H7 (ref. 4). Here we have sequenced the genome of E. coli O157:H7 to identify candidate genes responsible for pathogenesis, to develop better methods of strain detection and to advance our understanding of



**Figure 1** Circular genome map of EDL933 compared with MG1655. Outer circle shows the distribution of islands: shared co-linear backbone (blue); position of EDL933-specific sequences (O-islands) (red); MG1655-specific sequences (K-islands) (green); O-islands and K-islands at the same locations in the backbone (tan); hypervariable (purple). Second circle shows the G+C content calculated for each gene longer than 100 amino acids, plotted around the mean value for the whole genome, colour-coded like outer circle. Third

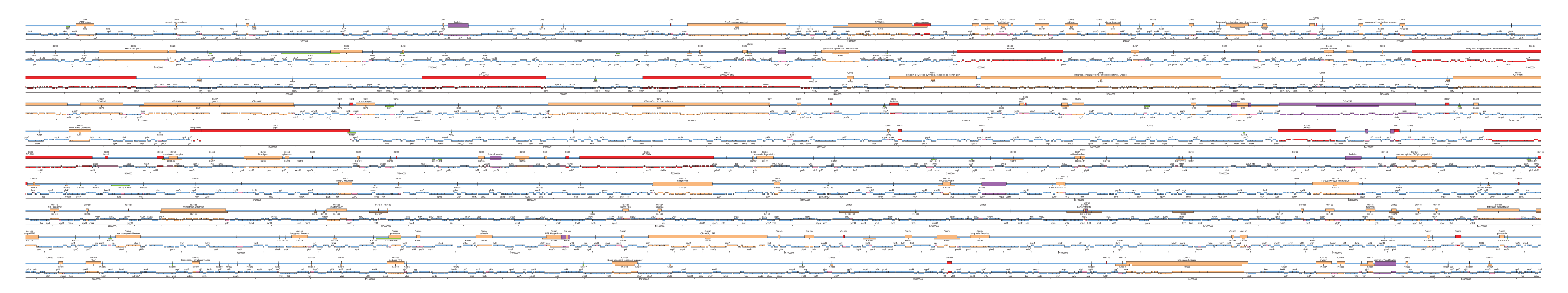
circle shows the GC skew for third-codon position, calculated for each gene longer than 100 amino acids: positive values, lime; negative values, dark green. Fourth circle gives the scale in base pairs. Fifth circle shows the distribution of the highly skewed octamer Chi (GCTGGTGG), where bright blue and purple indicate the two DNA strands. The origin and terminus of replication, the chromosomal inversion and the locations of the sequence gaps are indicated. Figure created by Genvision from DNASTAR.

the evolution of *E. coli*, through comparison with the genome of the non-pathogenic laboratory strain *E. coli* K-12 (ref. 5). We find that lateral gene transfer is far more extensive than previously anticipated. In fact, 1,387 new genes encoded in strain-specific clusters of diverse sizes were found in O157:H7. These include candidate virulence factors, alternative metabolic capacities, several prophages and other new functions—all of which could be targets for surveillance.

Escherichia coli O157:H7 was first associated with human disease after a multi-state outbreak in 1982 involving contaminated hamburgers<sup>1</sup>. The strain EDL933 that we sequenced was isolated from Michigan ground beef linked to this incident, and has been studied as a reference strain for O157:H7. Figures 1 and 2 show the gene content and organization of the EDL933 genome, and compare it with the chromosome of the K-12 laboratory strain MG1655 (ref. 5). These strains last shared a common ancestor about 4.5 million years ago<sup>6</sup>. The two E. coli genomes revealed an unexpectedly complex segmented relationship, even in a preliminary examination<sup>7</sup>. They share a common 'backbone' sequence which is co-linear except for one 422-kilobase (kb) inversion spanning the replication terminus (Fig. 1). Homology is punctuated by hundreds of islands of apparently introgressed DNA—numbered and designated 'K-islands' (KI) or 'O-islands' (OI) in Fig. 2, where K-islands are DNA segments present in MG1655 but not in EDL933, and Oislands are unique segments present in EDL933.

The backbone comprises 4.1 megabases (Mb), which are clearly homologous between the two *E. coli* genomes. O-islands total 1.34 Mb of DNA and K-islands total 0.53 Mb. These lineage-specific segments are found throughout both genomes in clusters of up to 88 kb. There are 177 O-islands and 234 K-islands greater than 50 bp in length. Histograms (Fig. 3) show more intermediate and large islands in EDL933 than in MG1655. Only 14.7% (26/177) of the O-islands correspond entirely to intergene regions. The two largest are identical copies of a 106-gene island, both in the same orientation and adjacent to genes encoding identical transfer RNAs.

Figure 2 Detailed comparative map of the EDL933 and MG1655 genomes. The upper double bar in each tier shows the genome comparison in EDL933 coordinates, with segments shown in detail and colour coded as in Fig. 1. Segments shown below the blue bar represent K-islands (MG1655-specific sequence). Segments extending above the blue backbone bar represent O-islands (EDL933-specific sequence). Unique identifying names (KI and OI numbers) were assigned to all segments of more than 50 bp. Unnamed vertical black lines across the blue bar indicate segments of less than 50 bp. In the lower line of each tier, EDL933 genes are presented showing orientation, and are coloured by segment type. Genes spanning segment junctions are shown in pink. Some gene names are given to provide landmarks in the backbone regions, and the sequence gaps are indicated. The scale in base pairs marks the base of each tier. Map created by Genvision from DNASTAR.



Labelling lineage-specific segments 'islands' is an extension of the term 'pathogenicity island' now in common, albeit varied, use. The original term arose from observations that virulence determinants are often clustered in large genomic segments showing hallmarks of horizontal transfer<sup>8</sup>. However, we found K- and O-islands of all sizes with no obvious association with pathogenicity; conversely, genes probably associated with virulence are not limited to the largest islands.

Roughly 26% of the EDL933 genes (1,387/5,416) lie completely within O-islands. In 189 cases, backbone-island junctions are within predicted genes. We classified the EDL933 genes into the functional groups reported for the MG1655 genome<sup>5</sup> and this is included in the annotation. Of the O-island genes, 40% (561) can be assigned a function. Another 338 EDL933 genes marked as unknowns lie within phage-related clusters and are probably remnants of phage genomes. About 33% (59/177) of the O-islands contain only genes of unknown function. Many classifiable proteins are related to known virulence-associated proteins from other *E. coli* strains or related enterobacteria.

Nine large O-islands (>15 kb) encode putative virulence factors: a macrophage toxin and ClpB-like chaperone (OI#7); a RTX-toxin-like exoprotein and transport system (OI#28); two urease gene clusters (OI#43 and #48); an adhesin and polyketide or fatty-acid biosynthesis system (OI#47); a type III secretion system and secreted proteins similar to the *Salmonella–Shigella inv–spa* host-cell invasion genes (OI#115); two toxins and a PagC-like virulence factor (OI#122); a fatty-acid biosynthesis system (OI#138); and the previously described locus of enterocyte effacement (OI#148). Among the large islands, four include a P4-family integrase and are directly adjacent to tRNAs (OI#43-serW, #48-serX, #122-pheV and #148-selC). Only the locus of enterocyte effacement and two of the lambdoid phages (see below) have as yet been experimentally associated with virulence in animal models.

Smaller islands that may be involved in virulence contain fimbrial biosynthesis systems, iron uptake and utilization clusters, and putative non-fimbrial adhesins. Many clusters have no obvious role in virulence, but may confer strain-specific abilities to survive in different niches. Examples include candidates for transporting diverse carbohydrates, antibiotic efflux, aromatic compound degradation, tellurite resistance and glutamate fermentation. Not all islands are expected to be adaptive. Some may represent neutral variation between strains. Still others may be deleterious but either have not yet been eliminated by selection or cannot be eliminated because of linkage constraints.

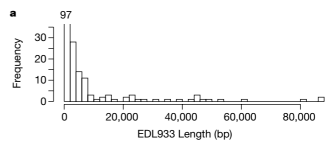
We identified 18 multigenic regions of the EDL933 chromosome related to known bacteriophages. Only one, the Stx2 Shiga toxinconverting phage BP-933W, is known to be capable of lytic growth and production of infectious particles<sup>10</sup>. We named the other EDL933 prophages cryptic prophage (CP) to indicate that they probably lack a full complement of functional phage genes. They vary in size from 7.5 kb (CP-933L) to 61.6 kb (BP-933W) and consist of a mosaic of segments similar to various bacteriophages, recalling the 'modular' phage genome hypothesis<sup>11</sup>. The two remaining physical gaps in the genome sequence correspond to prophage-related regions, and resolution of the sequence is complicated by extensive similarities to other prophage within this genome. The gap sizes and positions (4 kb and 54 kb) were determined from optical restriction maps. With only one exception, the EDL933 prophages and the eight cryptic prophages of MG1655 are all lineage-specific. Prophage Rac (MG1655) and CP-933R are similarly located in the backbone, and are sufficiently related to suggest a common prophage ancestor at the time that the strains

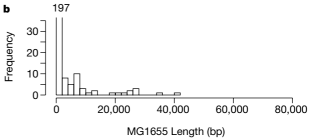
Subunits Stx1A and Stx1B of the second Shiga toxin of EDL933 (ref. 12) are encoded in the newly identified CP-933V. The position of the *stx1* genes in a putative Q antiterminator-dependent transcript is analogous to the placement of the *stx2* genes in BP-933W,

although there are no tRNAs adjacent to *stx1AB*. Genes in this position should be expressed maximally during lytic growth. The relationship between Stx toxin expression and phage induction is important, because treatment of O157:H7 with macrolide and quinolone antibiotics increase expression of the toxins<sup>13,14</sup>. Clinical decisions regarding drug therapy are complicated by strain-specific variation in this response<sup>15</sup>, and reports in the literature (for example, refs 6 and 12) taken together suggest that the Stx phage status is variable among O157:H7 strains. Given the potential for recombination among the prophage reported here, this does not seem surprising. In addition, the *stx* locus in *Shigella* is known to lie within a cryptic prophage, inserted at a site different from either *stx* phage of EDL933 (ref. 16).

The MG1655 genome contains 528 genes (528/4,405 = 12%) not found in EDL933. About 57% (303) of these were classified into known functional groups and include genes, such as for ferric citrate utilization, that would suggest a role in virulence if identified in a pathogen. It is unclear whether these are remnants of a recent pathogenic ancestor, steps along a path leading to evolution of a new pathogen, indicators that K-12 strains may be pathogenic for non-human hosts, or completely unrelated to pathogenicity. There are 106 examples of O-islands and K-islands present at the same locations relative to the conserved chromosomal backbone. The two replichores in each strain are nearly equal in length despite the large number of insertion/deletion events necessary to generate the observed segmented structure between strains. Only a subset of islands is associated with elements likely to be autonomously mobile.

Each island might be ancestral and lost from the reciprocal genome; however, atypical base composition suggests that most islands are horizontal transfers of relatively recent origin from a donor species with a different intrinsic base composition. Restricting analysis to the 108 O-islands greater than 1 kb, 94% (101/108) are significantly different ( $\chi^2 > 7.815$ , P < 0.05) from the average base composition of shared backbone regions in the same replichore. The percentage drops very little with a Bonferroni correction for multiple tests (91/108;  $\chi^2 > 17.892$ , P < 0.05). Similar results are obtained for analysis of the third-codon position composition (Fig. 1). Still more islands may have originated as horizontal transfers but have been resident in genomes with a spectrum of mutation similar enough to *E. coli* to have obtained equilibrium





**Figure 3** Histograms of lineage-specific segment lengths. **a**, EDL933; **b**, MG1655. The frequencies for the smallest length class are truncated to emphasize the distribution of longer clusters.

nucleotide frequencies or at least obscure statistical significance<sup>17</sup>. Still other gene clusters may be horizontal transfers that predate the divergence of MG1655 and EDL933.

Single nucleotide polymorphisms (75,168 differences) are distributed throughout the homologous backbone. There are 3,574 protein-coding genes encoded in backbone, and the average nucleotide identity for orthologous genes is 98.4%. Many orthologues (3,181/3,574 = 89%) are of equal length in the two genomes, but only 25% (911) encode identical proteins. Table 1 shows the number of each type of polymorphism observed by codon position. As expected, most differences are synonymous changes at third-codon positions. Multiple mutations at the same site should be infrequent at this low level of divergence. Thus the co-occurrence matrix provides insight into the substitution pattern, despite uncertainty of the ancestral state. The overall ratio of transitions to transversions is close to 3:1. A bias towards a greater number of T↔C than A↔G transitions on the coding strand previously attributed to transcription-coupled repair is evident<sup>18</sup>. An additional bias was observed at third-codon positions. Thymidines are more frequently involved in tranversions than cytosines, and G↔T are the most frequent transversions for the coding strand. The reciprocal polymorphisms,  $C \leftrightarrow A$ , are not over-represented. This bias is consistent for genes on both the leading and lagging strands (data not shown) and is therefore not related to asymmetries in the replication process. One possible explanation is transcription-coupled repair of damage associated with oxidative stress. Oxidized products of guanine (2,2,4-triaminooxazolone and 7,8-dihydro-8-oxoguanine) lead to G→T transversions by mispairing with A, and two DNA glycosylases (MutY and Fpg) are responsible for mismatch resolution<sup>19</sup>. Preferential repair of these lesions on the transcribed strand has been observed in humans<sup>20</sup>, and a similar mechanism could account for the observed transversion bias on the coding strand in E. coli.

Some chromosomal regions are more divergent ('hypervariable') than the average homologous segment but encode a comparable set of proteins at the same relative chromosomal position. In the most extreme case (YadC), the MG1655 and EDL933 proteins exhibit only 34% identity. Four such loci encode known or putative fimbrial

Table 1 Frequency of each type of single nucleotide polymorphism by codon position for 3,181 genes of equal length in EDL933 and MG1655

First-codon position					ED1 000 1 1 1
		Base in	MG1655		EDL933 totals
Base in EDL933 G A T C	G - 924 170 124 1,218	A 865 - 143 284 1,292	T 154 129 - 1,156 1,439	C 137 286 1,260 - 1,683	1,156 1,339 1,573 1,564 5,632
Second-codon posi	tion				
		Base in	MG1655		EDL933 totals
Base in EDL933 G A T C MG1655 totals	G - 393 67 107 567	A 410 - 147 176 733	T 66 159 - 394 619	C 118 166 464 - 748	594 718 678 677 2,667
Third-codon position	า				
		Base in	MG1655		EDL933 totals
Base in EDL933 G A T C MG1655 totals	G - 6,107 1,619 1,049 8,775	A 6,021 - 1,228 1,010 8,259	T 1,562 1,242 - 8,307 11,111	C 1,024 1,124 8,538 - 10,686	8,607 8,473 11,385 10,366 38,831

biosynthesis operons. Another encodes a restriction/modification system. Elevated divergence has been associated with positive selection at both these types of loci and among proteins that interact directly with the host<sup>9,21,22</sup>. Alternatively, hypervariable genes may result from locally elevated mutation rates or differential paralogue retention from an ancient tandem duplication.

Comparison of our observations with other genome-scale analyses of closely related strains or species supports the idea that enterobacterial genomes are particularly subject to recombinational evolution. Two Helicobacter pylori strains exhibited only 6-7% differential coding capacity despite showing less identity among orthologues (92.6%) than observed among these E. coli. Furthermore, almost half of the lineage-specific Helicobacter genes are clustered in a single region referred to as the plasticity zone<sup>23</sup>. Analyses of four Chlamydia genomes with orthologues that differ by as much as 19.5% show little evidence of horizontal transfer, and this is attributed to the inherent isolation of an obligate intracellular parasite<sup>24</sup>. Most lineage-specific genes are expansions of paralogous gene families. As in Helicobacter, many of the Chlamydia lineagespecific elements are clustered in a plasticity zone. Continuing genome projects will elucidate the generality of observations made from these comparisons of closely related organisms.

Together, our findings reveal a surprising level of diversity between two members of the species E. coli. Most differences in overall gene content are attributable to horizontal transfer, and offer a wealth of candidate genes that may be involved in pathogenesis. Base substitution has introduced variation into most gene products even among conserved regions of the two strains. Many of these differences can be exploited for development of highly sensitive diagnostic tools; but diagnostic utility will require a clearer understanding of the distribution of genetic elements in E. coli species as a whole. An independently isolated O157:H7 strain showed differences from EDL933 by restriction mapping<sup>25</sup>. Additional genome sequence data from other E. coli strains as well as functional characterization of gene products is necessary before the complex relationship between E. coli genotypes and phenotypes can be understood. Showing that disease-related traits are associated with predicted genes will require many areas of study including extensive testing in animal models that mimic symptoms of human infections, but the genome sequence offers a unique resource to help meet the challenge.

### Methods

### Clones and sequencing

EDL933 was kindly provided by C. Kaspar, who obtained it from the American Type Culture Collection (ATCC 43895). The sequenced isolate has been redeposited at the ATCC and is available as ATCC 700927. Whole-genome libraries in M13Janus and pBluescript were prepared from genomic DNA as described for genome segments used in the K-12 genome project<sup>26</sup>. Random clones were sequenced using dye-terminator chemistry and data were collected on ABI377 and 3700 automated sequencers. Sequence data were assembled by Seqman II (DNASTAR). Finishing used sequencing of opposite ends of linking clones, several PCR-based techniques and primer walking. Whole-genome optical maps for restriction enzymes NheI and XhoI were prepared<sup>27</sup> so that the ordering of contigs during assembly could be confirmed. Two gaps remain in the genome sequence. Extended exact matches pose a significant assembly challenge. The final determination of sequence for the 100-kb duplicated region was based on clones that span the junction between unique flanking sequences and the ends of the duplicated island, concordance of the two regions in optical restriction maps, excess random sequence coverage in the duplicated region, lack of polymorphism and confirmation of duplication of an internal segment by Southern blotting (data not shown).

#### Sequence features and database searches

Potential open reading frames (ORFs) were defined by GeneMark.hmm²8. The Gen-Pept118 protein and MG1655 protein and DNA databases were searched by each ORF using BLAST²9. Annotations were created from the search output in which each gene was inspected, assigned a unique identifier, and its product classified by functional group³5. Alternative start sites were chosen to conform to the annotated MG1655 sequence. Orthology was inferred when matches for EDL933 genes in the MG1655 database exceeded 90% nucleotide identity, alignments included at least 90% of both genes, and the MG1655 gene did not have an equivalent match elsewhere in the EDL933 genome. This list

was supplemented by manual inspection of the protein-level matches in the complete GenPept database to include genes with lower similarities if they occurred within co-linear regions of the genomes. The genome sequence was compared with that of MG1655 by the maximal exact match (MEM) alignment utility, (B.M., manuscript in preparation) an adaptation of MUMmer³0. This program was based on suffix arrays rather than suffix trees, and exact rather than unique matches, coupled with a custom anchored-alignment algorithm that extends sequence homology into the regions separating contiguous colinear exact matches. Inferences on biases in polymorphism patterns are based on  $\chi^2$  goodness-of-fit tests of a nested sequence of multinomial log–linear models. These predict symmetric elevated levels of A+G, T+C and G+T polymorphisms, above a quasi-independent baseline generated from marginal frequencies in the co-occurrence matrix of synonymous third-codon differences. Further information may be found at our Website http://www.genome.wisc.edu/, including a Genome Browser displaying a comparative map of EDL933 and K-12.

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# Genomic binding sites of the yeast cell-cycle transcription factors SBF and MBF

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Proteins interact with genomic DNA to bring the genome to life; and these interactions also define many functional features of the genome. SBF and MBF are sequence-specific transcription factors that activate gene expression during the G1/S transition of the cell cycle in yeast<sup>1,2</sup>. SBF is a heterodimer of Swi4 and Swi6, and MBF is a heterodimer of Mbp1 and Swi6 (refs 1, 3). The related Swi4 and Mbp1 proteins are the DNA-binding components of the respective factors, and Swi6 may have a regulatory function<sup>4,5</sup>. A small number of SBF and MBF target genes have been identified<sup>3,6-10</sup>. Here we define the genomic binding sites of the SBF and MBF transcription factors in vivo, by using DNA microarrays. In addition to the previously characterized targets, we have identified about 200 new putative targets. Our results support the hypothesis that SBF activated genes are predominantly involved in budding, and in membrane and cell-wall biosynthesis, whereas DNA replication and repair are the dominant functions among MBF activated genes<sup>6,11</sup>. The functional specialization of these factors may provide a mechanism for independent regulation of distinct molecular processes that normally occur in synchrony during the mitotic cell cycle.

To identify the targets of SBF and MBF, we combined chromatin immunoprecipitation and microarray hybridization (Fig. 1). Proteins were crosslinked with formaldehyde to their target sites *in vivo*. DNA that was specifically crosslinked to either of the transcription factors was purified by immunoprecipitation using an antibody against either the native protein or an epitope tag that was fused to the protein. Polymerase chain reaction (PCR) analysis of immunoprecipitated DNA confirmed the specific association of Swi4, Swi6 and Mbp1 with several known target promoters, and other target promoters that are identified here (see Supplementary Information). After reversal of the crosslinks, immunoprecipitated DNA was amplified and fluorescently labelled with the Cy5 fluoro-

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