1	The interpretation of temperature and salinity variables in numerical
2	ocean model output, and the calculation of heat fluxes and heat content
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4	by
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37 Abstract

The international thermodynamic equation of seawater of 2010 (TEOS-10) defined the enthalpy and entropy of seawater, thus enabling the global ocean heat content to be calculated as the volume integral of the product of in situ density, ρ , and potential enthalpy, h^0 (with reference sea pressure of 0 dbar). In terms of Conservative Temperature, Θ , ocean heat content is the volume integral of $\rho c_p^0 \Theta$, where c_p^0 is a constant "isobaric heat capacity".

44 However, many ocean models in the Coupled Model Intercomparison Project 45 phase 6 (CMIP6) as well as all models that contributed to earlier phases, such as 46 CMIP5, CMIP3, CMIP2 and CMIP1 used EOS-80 (Equation of State - 1980) rather than 47 the updated TEOS-10, so the question arises of how the salinity and temperature 48 variables in these models should be physically interpreted, with a particular focus on 49 comparison to TEOS-10 compliant observations. In this article we address how heat 50 content, surface heat fluxes and the meridional heat transport are best calculated using 51 output from these models, and how these quantities should be compared with those 52 calculated from corresponding observations. We conclude that even though a model 53 uses the EOS-80 equation of state which expects potential temperature as its input 54 temperature, the most appropriate interpretation of the model's temperature variable 55 is actually Conservative Temperature. This perhaps unexpected interpretation is 56 needed to ensure that the air-sea heat flux that leaves/arrives-in the atmosphere and 57 sea ice models is the same as that which arrives-in/leaves the ocean model.

We also show that the salinity variable carried by present TEOS-10 based models is Preformed Salinity, while the salinity variable of EOS-80 based models is also proportional to Preformed Salinity. These interpretations of the salinity and temperature variables in ocean models are an update on the comprehensive Griffies et al (2016) paper that discusses the interpretation of many aspects of coupled Earth system models.

65 **1. Introduction**

66 Numerical ocean models simulate the ocean by calculating the acceleration of 67 fluid parcels in response to various forces, some of which are related to spatially-varying 68 density fields that affect pressure, as well as solving transport equations for the two 69 tracers on which density depends, namely temperature [the CMIP6 variables identified 70 as thetao or bigthetao]) and dissolved matter ("salinity", [CMIP6 variable so]). For 71 computational reasons it is useful for the numerical schemes involved to be 72 conservative, meaning that the amount of heat and salt in the ocean changes only due to 73 the area integrated fluxes of heat and salt that cross the ocean's boundaries; in the case of 74 salt, this is zero. This conservative property is guaranteed for ocean models to within 75 computational truncation error since these numerical models are designed using finite 76 volume integrated tracer conservation (e.g., see Appendix F in Griffies et al 2016). It is 77 only by ensuring such conservation properties that scientists can reliably make use of 78 numerical ocean models for the long (centuries and longer) simulations required for 79 climate and Earth system studies.

However, this apparent numerical success ignores some difficult theoretical issues with the equation set being numerically solved. Here, we are concerned with issues related to the properties of seawater that have only recently been widely recognized because of research resulting in the Thermodynamic Equation of Seawater 2010 (TEOS-10). These issues mean that the intercomparison of different models, and comparison with ocean observations, needs to be undertaken with care.

86 In particular, it is widely recognized that the traditional measure of heat content 87 per unit mass in the ocean (with respect to an arbitrary reference state), the so-called 88 potential temperature, is not a conservative variable (McDougall, 2003). Hence, the time 89 change of potential temperature at a point in space is not determined solely by the 90 convergence of the potential temperature flux at that point. Furthermore, the non-91 conservative nature of potential temperature means that the potential temperature of a 92 mixture of water masses is not the mass average of the initial potential temperatures 93 since potential temperature is "produced" or "destroyed" by mixing within the ocean's 94 interior. This empirical fact is an inherent property of seawater (e.g., McDougall 2003,

Graham and McDougall 2013), and so treating potential temperature as a conservative
tracer (as well as making certain other assumptions related to the modelling of heat and
salt) results in contradictions, which have been built into most numerical ocean models
to varying degrees.

99 These contradictions have existed since the beginning of numerical ocean 100 modelling but have generally been ignored or overlooked because many other 101 oceanographic and numerical factors were of greater concern. However, as global heat 102 budgets and their imbalances are now a critical factor in understanding climate changes, 103 it is important to examine the consequences of these assumptions, and perhaps correct 104 them even at the cost of introducing problems elsewhere. These concerns are 105 particularly important when heat budgets are being compared between different 106 models, and with similar calculations made with observed conditions in the real ocean.

107 The purpose of this paper is to describe these theoretical difficulties, to estimate 108 the magnitude of errors that result, and to make recommendations about resolving them 109 both in current and future modelling efforts. For example, the insistence that a model's 110 temperature variable is potential temperature involves errors in the air-sea heat flux in 111 some areas that are as large as the mean rate of current global warming. A simple re-112 interpretation of the model's temperature variable overcomes this inconsistency and 113 allows coupled climate models to conserve heat.

114 The reader who wants to skip straight to the recommendations on how the 115 salinity and temperature outputs of CMIP models should be interpreted can go straight 116 to section 6.

117

118 **2. Background**

119 Thermodynamic measures of heat content

120 It is well-known that *in situ* temperature is not a satisfactory measure of the "heat 121 content" of a water parcel because the *in situ* temperature of a water parcel changes as 122 the ambient pressure changes (i.e., if a water parcel is transported to a different depth 123 [pressure] in the ocean). This change is of order 0.1°C as pressure changes 1000 dbar, 124 and is large relative to the precision of 0.01°C required to understand deep ocean circulation patterns. The utility of *in situ* temperature lies in the fact that it is easily
measured with a thermometer, and that air-sea boundary heat fluxes are to some degree
proportional to *in situ* temperature differences.

128 Traditionally, potential temperature has been used as an improved measure of 129 ocean heat content. Potential temperature is defined as the temperature that a parcel 130 would have if moved isentropically and without exchange of mass to a fixed reference 131 pressure (usually taken to be surface atmospheric pressure), and can be calculated from 132 measured ocean *in situ* temperatures using empirical correlation equations based on 133 laboratory measurements. However, the enthalpy of seawater varies nonlinearly with 134 temperature and salinity (Fig. 1) and this variation results in non-conservative behaviour 135 under mixing (McDougall (2003), section A.17 of IOC et al. (2010)). The ocean's potential 136 temperature is subject to internal sources and sinks – it is not conservative.

With the development of a Gibbs function for seawater, based on empirical fits to measurements of known thermodynamic properties (Feistel (2008), IOC et al, 2010), it became possible to apply a more rigorous theory for quasi-equilibrium thermodynamics to study heat content problems in the ocean. As a practical matter, calculations can now be made that allow for an estimate of the magnitude of non-conservative terms in the ocean circulation. By integrating over water depth these production rates can be expressed as an equivalent heat flux per unit area.

144 Non-conservation of potential temperature was found to be equivalent to a root 145 mean square surface heat flux of about 60 mWm⁻² (Graham and McDougall, 2013), and 146 an average value of 16 mWm⁻² (see below). These numbers can be compared to a 147 present-day estimated global-warming surface heat flux imbalance of between 148 300 mWm^{-2} and 470 mWm^{-2} (Zanna et al., 2019, von Schuckmann et al., 2020). By 149 comparison, the globally averaged rate of increase of temperature due to the dissipation 150 of kinetic energy is approximately 10 mW m⁻². These equivalent heat fluxes and 151 subsequent similar values are gathered into Table 1 for reference. In the context of a 152 conceptual ocean model being driven by known heat fluxes, the presence of the non-153 conservation of potential temperature causes SST errors seasonally in the equatorial 154 region of about 0.5K (0.5°C), while the error (in all seasons) at the outflow of the

Amazon is 1.8K (see section 9 of McDougall, 2003). With different boundary conditions (such as restoring boundary conditions) the error in assuming that potential temperature is conservative is split in different proportions, between (a) the potential temperature values and (b) the potential temperature fluxes.

Unfortunately, no single alternative thermodynamic variable has been found that is both independent of pressure, and conservative under mixing. For example, specific entropy is produced in the ocean interior when mixing occurs, with the depth-integrated production being equivalent to an imbalance in the air-sea heat flux of a root mean square value of about 500 mW m^{-2} (Graham and McDougall, 2013), while, apart from the dissipation of kinetic energy, enthalpy is conservative under mixing at constant pressure, but enthalpy is intrinsically pressure-dependent.

166 However, it was found that a constructed variable, potential enthalpy 167 (McDougall, 2003), has a mean non-conservation error in the global ocean of only about 168 0.3 mW m^{-2} (this is the mean value of an equivalent surface heat flux, equal to the depth 169 integrated interior production of potential enthalpy that is additional to the production 170 due to the dissipation of kinetic energy (Graham and McDougall, 2013)). The potential 171 enthalpy, \tilde{h}^0 , is the enthalpy of a water parcel after being moved adiabatically and at 172 constant salinity to the reference pressure 0 dbar where the temperature is equal to the 173 potential temperature, θ , of the water parcel:

174

$$\tilde{h}^0(S_{\scriptscriptstyle A},\theta) = h(S_{\scriptscriptstyle A},\theta,0\,\text{dbar}). \tag{1}$$

175 In Eq. (1) the function h is the specific enthalpy of TEOS-10 (defined as a function of Absolute Salinity, in-situ temperature and sea pressure) whereas \tilde{h}^0 is the potential 176 177 enthalpy function and the over-twiddle implies that the temperature input to this 178 function is potential temperature, θ . By way of comparison, the area-averaged 179 geothermal input of heat into the ocean bottom is about 86 mWm⁻², and the interior 180 heating of the ocean due to viscous dissipation, is equivalent to a mean surface heat flux 181 of about 3 mWm⁻² (Graham and McDougall, 2013). Tailleux (personal communication, 182 2021) has suggested that the dissipation of kinetic energy in the ocean may be as much 183 as three times as large as this value, at approximately 10 mW m⁻². Thus we conclude 184 that potential enthalpy, although not a theoretically ideal conservative parameter, can be

185 treated as such for many present purposes in oceanography. If at some stage in the 186 future a source term were to be added to the evolution equation for Conservative 187 Temperature, the most important contribution would be that due to the dissipation of 188 kinetic energy, being a factor of ~10-30 larger than the non-conservation of Conservative 189 Temperature due to other diffusive contributions (namely the terms on the last two lines 190 of Eqn. (38) of Graham and McDougall (2013)).

191 Since potential enthalpy was not a widely understood property, a decision was
192 made in the development of TEOS-10 to adopt Conservative Temperature, Θ, which has
193 units of temperature and is proportional to potential enthalpy:

194

$$\Theta = \tilde{\Theta}(S_{A}, \theta) = \tilde{h}^{0}(S_{A}, \theta) / c_{p}^{0}, \qquad (2)$$

195 where the proportionality constant $c_p^0 \equiv 3991.867\,957\,119\,63\,\,\mathrm{Jkg}^{-1}\mathrm{K}^{-1}$, was chosen so that 196 the average value of Conservative Temperature at the ocean surface matched that of 197 potential temperature. Although in hindsight other choices (e.g., with fewer significant 198 digits) might have been more useful, this value of c_p^0 is now built into the TEOS-10 199 standard.

Note that at specific locations in the ocean, in particular at low salinities and high temperatures, Θ and θ can differ by more than 1°C (Fig. 2); the difference is a strongly nonlinear function of temperature and salinity. Θ is, by definition, independent of adiabatic and isohaline changes in pressure.

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205 Why is potential temperature not conservative?

206 This question is answered in sections A.17 and A.18 of the TEOS-10 Manual (IOC 207 et al., 2010) as well as McDougall (2003) and Graham and McDougall (2013). The answer 208 is that potential enthalpy referenced to the sea surface pressure, h^0 , which is an (almost 209 totally) conservative variable in the real ocean, is not simply a linear function of 210 potential temperature, θ , and Absolute Salinity, S_{A} (and note that both enthalpy and 211 entropy are unknown and unknowable up to separate linear functions of Absolute 212 Salinity). If potential enthalpy were a linear function of potential temperature and 213 Absolute Salinity then the "heat content" per unit mass of seawater could be accurately 214 taken to be proportional to potential temperature, and the isobaric specific heat capacity

at zero sea pressure would be a constant. As an example of the nonlinearity of $\tilde{h}^0(S_A, \theta)$, the isobaric specific heat at the sea surface pressure $c_p(S_A, \theta, 0 \text{ dbar}) \equiv h_{\theta}^0$ varies by 6% across the full range of temperatures and salinities found in the World Ocean (Fig. 1). By way of contrast, the potential enthalpy of an ideal gas is proportional to its potential temperature.

Another way of treating heat in an ocean model is to continue carrying potential temperature as its temperature variable but to (i) use the variable isobaric heat capacity at the sea surface to relate the air-sea heat flux to an air-sea flux of potential temperature, and (ii) to evaluate the non-conservative source terms of potential temperature and add these source terms to the potential temperature evolution equation during the ocean model simulation (Tailleux, 2015).

226 However it is not possible to accurately choose the value of the isobaric heat 227 capacity at the sea surface that is needed when θ is the model's temperature variable. This 228 issue arises because of the unresolved variations in the sea surface salinity (SSS) and SST (for 229 example, unresolved rain events that temporarily lower the SSS), together with the nonlinear 230 dependence of the isobaric specific heat on salinity and temperature. Because of such 231 unresolved correlations, the air-sea heat flux would be systematically mis-estimated. 232 Neither is it possible to accurately estimate the non-conservative source terms of θ in the 233 ocean interior. This problem arises because the source terms are the product of a turbulent 234 flux and a mean gradient. In a mesoscale eddy-resolved ocean model (or even finer scale) it 235 is not clear how to find the eddy flux of θ , as this depends on how the averaging is done in 236 space and time. Furthermore, when analysing the output of such an ocean model, one 237 would need to find a way of dealing with the contributions from source terms that are not 238 expressible in the form of flux convergences when, for example, estimating the meridional 239 heat transport.

We conclude that the idea that ocean models could retain potential temperature θ as the model's temperature variable, rather than adopt the TEOS-10 recommendation of using Conservative Temperature Θ , is unworkable because (1) the air-sea heat flux cannot be accurately evaluated, (2), the non-conservative source terms that appear in the θ evolution equation cannot be estimated accurately, and (3) the ocean section-integrated heat fluxescannot be accurately calculated.

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7 How conservative is Conservative Temperature?

248 This question is addressed in McDougall (2003) as well as in section A.18 of the 249 TEOS-10 Manual (IOC et al., 2010) and in Graham and McDougall (2013). The first step 250 in addressing the non-conservation of Θ is to find a thermophysical variable that is 251 conserved when fluid parcels mix. McDougall (2003) and Graham and McDougall 252 (2013) showed that when fluid parcels are brought together adiabatically and 253 isentropically to mix at pressure p^m , it is the potential enthalpy h^m referenced to the 254 pressure p^m of a mixing event that is conserved, apart from the dissipation of kinetic 255 From this knowledge they constructed the evolution equations for energy, ε . 256 Conservative Temperature as well as for potential temperature and for entropy.

257 By contrast, Tailleux (2010) and Tailleux (2015) assumed that it was the Total 258 Energy, being the sum of internal energy, kinetic energy and the geopotential, that is 259 conserved when fluid parcels mix in the ocean. However, as shown by McDougall, 260 Church and Jackett (2003), the $-\nabla \cdot (P\mathbf{u})$ term on the right-hand side of the evolution 261 equation for Total Energy is non-zero when integrated over the mixing region, so that 262 Total Energy is not a conservative variable. Tailleux (2010, 2015) treated this nonconservative term, $-\nabla \cdot (P\mathbf{u})$, as though it were a conservative term in all their evolution 263 264 equations, so that these papers actually arrived at the correct evolution equations for 265 Θ , θ and η (for example, Eqn. (B.7) of Tailleux (2010) and Eqn. (B10) of Graham and 266 McDougall (2013) are identical). However, these equations are written in terms of the 267 molecular fluxes of heat and salt, and the Tailleux (2010, 2015) papers did not find a way 268 to use these expressions to evaluate the non-conservation of Θ , θ and η in a turbulently 269 mixed ocean. This was done in section 3 of Graham and McDougall (2013).

While enthalpy is conserved when mixing occurs at constant pressure, it does not possess the "potential" property, but rather, an adiabatic and isohaline change in pressure causes a change in enthalpy according to $\hat{h}_p = v$, where v is the specific volume. This property is illustrated in Fig. 3 where it is seen that for an adiabatic and 274 isohaline increase of pressure of 1000dbar, the increase in enthalpy is the same as that 275 caused by an increase in Conservative Temperature of more than 2.4°C. If enthalpy 276 variations at constant pressure were a linear function of Absolute Salinity and 277 Conservative Temperature, the contours in Fig. 3 would be parallel equidistant straight 278 lines, and Conservative Temperature would be totally conservative. Since this is not the 279 case, this figure illustrates the (small) non-conservation of Conservative Temperature. 280 Further discussion and evaluation of the non-conservation of Conservative Temperature 281 can be found in McDougall (2003) and Graham and McDougall (2013).

282

283 Seawater Salinity

284 To a degree of approximation which is useful for many purposes, the dissolved 285 matter in seawater ("sea salt") can be treated as a material of uniform composition, 286 whose globally averaged absolute salinity (i.e. the grams of solute per kilogram of 287 seawater) changes only due to the addition and removal of freshwater through rain, 288 evaporation, and river inflow. This property is because the processes that govern the 289 addition and removal of the constituents of sea salt have extremely long time scales, 290 relative to those that affect the pure water component of seawater. We can thus treat the 291 total ocean salt content as approximately constant, while subject to spatially and 292 temporally varying boundary fluxes of fresh water that give rise to salinity gradients.

293 The utility of this definition of uniform composition of sea salt lies in its 294 conceptual simplicity, well suited to theoretical and numerical ocean modelling at time 295 scales of up to 100s of years. However, to the demanding degree required for observing 296 and understanding deep ocean pressure gradients, sea salt is neither uniform in 297 composition, nor is it a conserved variable, nor can its absolute amount be measured 298 precisely in practice. The repeatable precision of various technologies used to estimate 299 salinity can be as small as 0.002 g/kg, but the non-ideal nature of seawater means that 300 these estimates can be different by as much as 0.025 g/kg relative to the true Absolute 301 Salinity in the open ocean, and as much 0.1 g/kg in coastal areas (Pawlowicz, 2015).

The most important interior source and sink factors governing changes in the composition of sea salt are biogeochemical processes that govern the biological uptake of dissolved nutrients, calcium, and carbon in the upper ocean, and the remineralization of these substances from sinking particles at depth. At present it is thought that changes resulting from hydrothermal vent activity, fractionation from sea ice formation, and through multi-component molecular diffusion processes are of local importance only, but little work has been done to quantify this.

To address this problem, TEOS-10 defines a Reference Composition of seawater, and several slightly different salinity variables that are necessary for different purposes to account for the variable composition of sea salt. The TEOS-10 Absolute Salinity, S_A , is the absolute salinity of Reference Composition Seawater of a measured density (note that capitalization of variable names denotes a precise definition in TEOS-10). It is the salinity variable that is designed to be used to accurately calculate density using the TEOS-10 Gibbs function.

316 Preformed Salinity, S_* , is the salinity of a seawater parcel with the effects of 317 biogeochemical processes removed, somewhat analogous to a chlorinity-based salinity 318 estimate. It is thus a conservative tracer of seawater, suitable for modelling purposes, 319 but neglects the spatially variable small portion of sea salt involved in biogeochemical 320 processes that is required for the most accurate density estimates. Since the original 321 measurements of specific volume to which both EOS-80 and TEOS-10 were fitted were 322 made on samples of Standard Seawater with composition close to Reference 323 Composition, the Reference Salinity of these samples were also the same as Preformed 324 Salinity.

325 Ocean observational databases contain a completely different variable; Practical 326 Salinity. This variable, which predates TEOS-10, is essentially based on a measure of the 327 electrical conductance of seawater, normalized to conditions of fixed temperature and 328 pressure by empirical correlation equations, between the ranges of 2 and 42 PSS-78 and 329 scaled so that ocean salinity measurements that have been made through a variety of 330 technologies over the past 120 years are numerically comparable. Practical Salinity 331 measurement technologies involve a certified reference material called IAPSO Standard 332 Seawater, which for our purposes can be considered the best available artifact 333 representing seawater of Reference Composition.

Practical Salinity was not designed for numerical modelling purposes and does not accurately represent the mass fraction of dissolved matter. We can link Practical Salinity, $S_{\rm p}$, to the Absolute Salinity of Reference Composition seawater (so-called Reference Salinity, $S_{\rm R}$) using a fixed scale factor, $u_{\rm PS}$, so that

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$$S_{\rm R} = u_{\rm PS} S_{\rm P}$$
 where $u_{\rm PS} \equiv (35.165\ 04/35)\ {\rm gkg}^{-1}$. (3)

339 Conversions to and between the other "salinity" definitions, however, involve 340 knowledge about spatial and temporal variations in seawater composition. Fortunately, 341 the largest component of these changes occurs in a set of constituents involved in 342 biogeochemical processes, whose co-variation is known to be strongly correlated. Thus 343 the Absolute Salinity of real seawater can be determined globally to useful accuracy 344 from the Reference Salinity by the addition of a single parameter, the so-called Absolute 345 Salinity Anomaly, δS_A ,

$$S_{\rm A} = S_{\rm R} + \delta S_{\rm A} \,, \tag{4}$$

which has been tabulated in a global atlas for the current ocean (McDougall et al., 2012),
and is estimated in coastal areas by considering the effects of river salts (Pawlowicz,
2015). It can also be determined from measurements of either density or of carbon and
nutrients (IOC et al., 2010, Ji et al., 2021). For purposes of numerical ocean modelling,
the Absolute Salinity Anomaly could in theory be obtained by separately tracking the
carbon cycle and nutrients, and applying known correction factors, but we are not aware
of any attempts to do so.

354 Chemical modelling (Pawlowicz (2010), Wright et al. (2011), Pawlowicz et al.355 (2012)) suggests the approximate relation

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$$S_{\rm A} - S_* \approx 1.35 \,\delta S_{\rm A} \equiv 1.35 (S_{\rm A} - S_{\rm R}),$$
 (5)

and these relationships are schematically illustrated in Fig. 4. The magnitude of the Absolute Salinity Anomaly is around -.005 to +0.025 g/kg in the open ocean, relative to a mean Absolute Salinity of about 35 g/kg. The correction it implies may be important when initializing models, or comparing them with observations, but its major effect is likely in producing biases in calculated isobaric density gradients.

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365 The density of seawater is the most important thermodynamic property affecting 366 oceanic motions, since its spatial changes (along with changes to the sea-surface height) 367 give rise to pressure gradients which are the primary driving force for currents within 368 the ocean interior through the hydrostatic relation. The "traditional" equation of state is 369 known as EOS-80 (UNESCO, 1981), and is standardized as a function of Practical 370 Salinity and in-situ temperature, $\rho = \rho(S_{\rm p}, t, p)$ which has 41 numerical terms. An 371 additional equation (the adiabatic lapse rate) is required for conversion of temperature 372 to potential temperature. However, for ocean models, the EOS-80 equation of state is 373 usually taken to be the 41-term expression written in terms of potential temperature, 374 $\rho = \tilde{\rho}(S_{\rm p}, \theta, p)$, of Jackett and McDougall (1995), where the over-twiddle indicates that 375 this rational function fit was made with Practical Salinity S_{p} and potential temperature

376 θ as the input salinity and temperature variables.

The current standard for describing the thermodynamic properties of seawater, known as TEOS-10, provides an equation of state, $v = 1/\rho = v(S_A, t, p)$, in the form of a function which involves 72 coefficients (IOC et al., 2010) and is an analytical pressure derivative of the TEOS-10 Gibbs function. However, for ocean models using TEOS-10 the equation of state used is one of those in Roquet et al. (2015); the 55-term equation of state, $\rho = \hat{\rho}(S_A, \Theta, z)$, used by Boussinesq models and the 75-term polynomial for specific volume, $v = \hat{v}(S_A, \Theta, p)$, used by non-Boussinesq ocean models.

In this paper we will not concentrate on the distinction between Boussinesq and non-Boussinesq ocean models, and henceforth we will take the third input to the equation of state to be pressure, even though for a Boussinesq model it is in fact a scaled version of depth as per the energetic arguments of Young (2010). By the same token, we will cast the discussion in terms of the *in situ* density, even though the non-Boussinesq models have as their equation of state a polynomial for the specific volume, $v = 1/\rho$.

For seawater of Reference Composition, both the TEOS-10 and EOS-80 fits $\rho = \hat{\rho}(S_A, \Theta, p)$ and $\rho = \tilde{\rho}(S_P, \theta, p)$ are almost equally accurate (see section A.5 of IOC et al. (2010) and note the comparison between Figures A.5.1 and A.5.2 therein). That is, if we set $\delta S_A = 0$ and use Eqn. (3) to relate Practical and Reference Salinities (which in this case are the same as Preformed Salinities), the numerical density values of in situ density
calculated using EOS-80 are not significantly different to those using TEOS-10 in the
open ocean [the differences are significant for brackish waters].

397 This being the case, we can see from sections A.5 and A.20 of the TEOS-10 398 Manual (IOC et al. (2010)) that 58% of the data deeper than 1000 dbar in the World 399 Ocean would have the thermal wind misestimated by ~2.7% due to ignoring the 400 difference between Absolute and Reference Salinities. No ocean model has addressed 401 this deficiency to date, but McCarthy et al. (2015) studied the influence of using Absolute 402 Salinity versus Reference Salinity in calculating the overturning circulation in the North 403 Atlantic. They found that the overturning streamfunction changed by 0.7Sv at a depth 404 of 2700m, relative to a mean value at this depth of about 7 Sv, i.e., a 10% effect. Because 405 we argue that the salinity variable in ocean models is best interpreted as being 406 Preformed Salinity, S_* , the neglect of the distinction between Preformed and Absolute 407 Salinities in ocean models means that they misestimate the overturning streamfunction 408 by 1.35 (see Figure 4) times 0.7Sv, namely ~1Sv, i.e., a 13.5% effect.

409

410 Air-sea heat fluxes

411 Sensible, latent and long-wave radiative fluxes are affected by near-surface 412 turbulence and are usually calculated using bulk formulae involving air and sea 413 surface water temperatures (the air and sea *in situ* temperatures), as well as other 414 parameters (e.g., the latent heat involves the isobaric evaporation enthalpy, commonly 415 called the latent heat of evaporation, which is actually a weak function of temperature 416 and salinity; see Eqn. 6.28 of Feistel et al. (2010) and Eqn. (3.39.7) of IOC et al. (2010)). 417 The total air-sea heat flux, *Q*, is then translated into a water temperature change by 418 dividing by a heat capacity c_n^0 , which has always been taken to be constant in 419 numerical models (Griffies et al., 2016). Although this method is appropriate for Conservative Temperature, CT, (assuming that the TEOS-10 value is used for c_p^0), it is 420 421 not appropriate when potential temperature is being considered. The flux of potential 422 temperature into the surface of the ocean should be Q divided by $c_n(S_*, \theta, 0)$. The use 423 of a constant specific heat capacity, in association with the interpretation of the

424 ocean's temperature variable as being potential temperature, means that the ocean has
425 received a different amount of heat than the atmosphere actually delivers to the ocean,
426 and this issue will be explored in section 3.

When precipitation (*P*) occurs at the sea surface, this addition of freshwater brings with it the associated potential enthalpy $h(S_A = 0, t, 0 \text{ dbar})$ per unit mass of freshwater, where *t* is the *in situ* temperature of the rain drops as they arrive at the sea surface. The temperature at which rain enters the ocean is not yet treated consistently in coupled models, and section K1.6 of Griffies et al. (2016) suggests that this effect could be equivalent to an area-averaged extra air-sea heat flux of between -150 mW m^{-2} and -300 mW m^{-2} , representing a heat loss for the ocean.

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436 Numerical ocean models

In deciding how to numerically model the ocean, an explicit choice must be made about the equation of state, and one would think that this choice would have implications about the precise meaning of the temperature and salinity variables in the model, which we will call T_{model} and S_{model} respectively. We can divide ocean models into two general classes, EOS-80 models, and TEOS-10 models:

442

443 <u>EOS-80 models</u>

444 One class of CMIP ocean model is based around EOS-80, and these models have the445 following characteristics:

- 446 1. The model's equation of state, $\rho = \tilde{\rho}(S_p, \theta, p)$, expects to have Practical Salinity 447 and potential temperature as the salinity and temperature input parameters.
- 448
 2. T_{model} is advected and diffused in the ocean interior in a conservative manner, i.e.,
 449 its evolution at a point in space is determined by the convergence of advective
 450 fluxes plus parameterized sub-grid scale diffusive and skew diffusive fluxes.
- 451 3. S_{model} is advected and diffused in the ocean interior in a conservative manner as 452 for T_{model} .

- 453 4. The air-sea heat flux is delivered to/from the ocean using a constant isobaric specific heat, c_n^0 , to convert the air-sea heat flux into a surface flux of T_{model} . [An 454 EOS-80 based model's value of c_p^0 is generally only slightly different to the 455 456 TEOS-10 value.]
- 457

5. T_{model} is initialized from an atlas of values of potential temperature, and S_{model} is 458 initialized with values of Practical Salinity.

459 At first glance, it seems reasonable to assume that T_{model} is potential temperature, and $S_{\rm model}$ is Practical Salinity. However, these assumptions imply that theoretical errors 460 461 arising from items 2 and 3 and 4 are ignored (since neither potential temperature nor 462 Practical Salinity are conservative variables). In this paper we show that these 463 interpretations of the model's temperature and salinity variables are not as accurate as 464 our proposed alternative interpretations.

465

466 TEOS-10 models

467 Other ocean models have begun to implement TEOS-10 features. These models 468 generally have the following characteristics.

- 469 1. The model's equation of state, $\rho = \hat{\rho}(S_A, \Theta, p)$, expects to have Absolute Salinity 470 and Conservative Temperature as its salinity and temperature input parameters.
- 471 2. T_{model} is advected and diffused in the ocean interior in a conservative manner.
- $S_{\rm model}$ is advected and diffused in the ocean interior in a conservative manner. 472 3.
- 473 4. At each time step of the model, the value of potential temperature at the sea 474 surface (i.e. SST) is calculated from the T_{model} (which is assumed to be 475 Conservative Temperature) and this value of SST is used to interact with the 476 atmosphere via bulk flux formulae.
- 477 5. The air-sea heat flux is delivered to/from the ocean using the TEOS-10 constant 478 isobaric specific heat, c_p^0 , to convert the air-sea heat flux into a surface flux of 479 T_{model} .
- 6. T_{model} is initialized from an atlas of values of Conservative Temperature, and 480 481 $S_{\rm model}$ is initialized with values of one of Absolute Salinity, Reference Salinity or 482 Preformed Salinity.

483 Implicitly, it has then been assumed that T_{model} is a Conservative Temperature, and S_{model} 484 is Absolute Salinity.

There is one CMIP6 ocean model that we are aware of, ACCESS-CM2 (Australian Community Climate and Earth System Simulator, Bi et al. 2013), whose equation of state is written in terms of Conservative Temperature, but the salinity argument in the equation of state is Practical Salinity. The salinity in this model is initialized with atlas values of Practical Salinity.

From the above it is clear that there are small but significant theoretical incompatibilities between different models, and between models and the observed ocean. These issues become apparent when dealing with the technicalities of intercomparisons, and various choices must be made. We now consider the implications of these different choices and provide recommendations for best practices.

495

496 **3.** The Interpretation of salinity in ocean models

497 Note that the samples whose measured specific volumes were incorporated into 498 both the EOS-80 and TEOS-10 equations of state were of Standard Seawater whose 499 composition is close to Reference Composition. Consequently, the EOS-80 and TEOS-10 500 equations of state were constructed with Preformed Salinity, S_* (or, in the case of EOS-501 80 models, $S_*/u_{\rm PS}$), as their salinity arguments, not Reference Salinity. These same 502 algorithms give accurate values of specific volume for seawater samples that are not of 503 Reference Composition so long as the salinity argument is Absolute Salinity (as opposed 504 to Reference Salinity or Preformed Salinity).

505 For an ocean model that has no non-conservative interior source terms affecting 506 the evolution of its salinity variable, and that is initialized at the sea surface with 507 Preformed Salinity, the only interpretation for the model's salinity variable is Preformed 508 Salinity, and the use of the TEOS-10 equation of state will then yield the correct specific 509 volume. Furthermore, whether the model is initialized with values of Absolute Salinity, 510 Reference Salinity or Preformed Salinity, these initial salinity values are nearly identical 511 in the upper ocean, and any differences between the three initial conditions in the 512 deeper ocean would be largely diffused away within the long spin-up period. That is, in

the absence of the non-conservative biogeochemical source terms that would be needed to model Absolute Salinity and to force it away from being conservative (or the smaller source terms that would be needed to maintain Reference Salinity), the model's salinity variable will drift towards being Preformed Salinity. Hence, we conclude that, after the long spin-up phase, the salinity variable of a TEOS-10 based ocean model is accurately interpreted as being Preformed Salinity S_* , irrespective of whether the model was initialized with values of Absolute Salinity, Reference Salinity or Preformed Salinity.

520 Likewise, the prognostic salinity variable after a long spin-up period of an EOS-521 80 based model is most accurately interpreted as being Preformed Salinity divided by 522 $u_{\rm PS} \equiv (35.16504/35) \, {\rm gkg}^{-1}$, $S_*/u_{\rm PS}$.

523 We clearly need more estimates of the magnitude of the dynamic effects of the 524 variable seawater composition, but for now we might take a change in 1 Sv in the 525 meridional transport of deep water masses in each ocean basin (based on the Atlantic 526 work of McCarthy et al., 2015) as an indication of the magnitude of the effect of 527 neglecting the effects of biogeochemistry on salinity. At this stage of model 528 development, since all models are equally deficient in their thermophysical treatment of 529 salinity, at least this aspect does not present a problem as far as making comparisons 530 between CMIP models.

531

532 4. Model Heat Flux Calculations

533 From the details described above, both types of numerical ocean models suffer from 534 some internal contradictions with thermodynamical best practice. For example, for the 535 EOS-80 based models, if T_{model} is assumed to be potential temperature, the use of EOS-80 is correct for density calculations but the use of conservative equations for T_{model} ignores 536 537 the non-conservative production of potential temperature. The use of a constant heat 538 capacity is also in error if T_{model} is interpreted as potential temperature. Conservative 539 equations are, however, appropriate for Conservative Temperature. In addition, if S_{model} 540 is assumed to be either Practical Salinity or Absolute Salinity, then the use of 541 conservative equations ignores the changes in salinity that arise from biogeochemical 542 processes.

543 One use for these models is to calculate heat budgets and heat fluxes – both at the 544 surface and between latitudinal bands, and inherent to CMIP is the idea that these 545 different models should be intercompared. The question of how this intercomparison 546 should be done, however, was not clearly addressed in Griffies et al. (2016). Here we 547 begin the discussion by considering two different options for interpreting T_{model} in EOS-548 80 ocean models.

549

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553

550 4.1 Option 1: interpreting the EOS-80 model's temperature as being potential

551

temperature

Under this option the model's temperature variable T_{model} is treated as being potential temperature θ ; this is the prevailing interpretation to date. With this interpretation of

554 T_{model} one wonders whether Conservative Temperature Θ should be calculated from the 555 model's (assumed) potential temperature before calculating (i) the global Ocean Heat 556 Content as the volume integral of $\rho c_n^0 \Theta$, and (ii) the advective meridional heat transport 557 as the area integral of $\rho c_n^0 \Theta v$ at constant latitude, where v is the northward velocity. 558 This question was not clearly addressed in Griffies et al. (2016), and here we emphasize 559 one of the main conclusions of the present paper, namely that ocean heat content and 560 meridional heat transports should be calculated using the model's prognostic 561 temperature variable. Any subsequent conversion from one temperature variable to 562 another (such as potential to Conservative) in order to calculate heat content and heat 563 transport is incorrect and confusing, and should not be attempted.

564

565 4.1.1 Issues with the potential temperature interpretation

566 There are several thermodynamic inconsistencies that arise from option 1. First, the ocean model has assumed in its spin-up phase (for perhaps a millennium) that T_{model} 567 568 is conservative, so during the whole spin-up phase and beyond, the contribution of the 569 known non-conservative interior source terms of potential temperature have been 570 absent, and hence the model's temperature variable has not responded to these absent 571 source terms and so this temperature field cannot be potential temperature. Also, since 572 the temperature field of the model is not potential temperature (because of these absent

source terms) the velocity field of the model will also not be forced correctly due toerrors in the density field which in turn affect the pressure force.

575 The second inconsistent aspect of option 1 is that the air-sea flux of heat is 576 ingested into the ocean model, both during the spin-up stage and during the subsequent 577 transient response phase, as though the model's temperature variable is proportional to 578 potential enthalpy. For example, consider some time during the year at a particular 579 location where the sea surface is fresh (a river outflow, or melted ice). During this time, 580 any heat that the atmosphere loses or gains should have affected the potential 581 temperature of the upper layers of the ocean using a specific heat that is 6% larger than 582 c_n^0 (see Figure 1). So, if the ocean model's temperature variable is interpreted as being 583 potential temperature, a 6% error is made in the heat flux that is exchanged with the 584 atmosphere during these periods/locations. That is, the changes in the ocean model's 585 (assumed) potential temperature caused by the air-sea heat flux will be exaggerated 586 where and when the sea surface salinity is fresh. This 6% flux error is not corrected by 587 subsequently calculating Conservative Temperature from potential temperature; for 588 example, these temperatures are the same at low temperature and salinity (see Figure 2), 589 and yet at low values of salinity, the specific heat is 6% larger than c_n^0 .

590 This second inconsistent aspect of option 1 can be restated as follows. The 591 adoption of potential temperature as the model's temperature variable means that there 592 is a discontinuity in the heat flux of the coupled air-sea system right at the sea surface; 593 for every Joule of heat (i.e. potential enthalpy) that the atmosphere gives to the ocean, 594 under this Option 1 interpretation, up to 6% too much heat arrives in the ocean over 595 relatively fresh waters. In this way, the adoption of potential temperature as the model 596 temperature variable ensures that the coupled ocean atmosphere system will not 597 conserve heat. Rather, there appear to be non-conservative sources and sinks of heat 598 right at the sea surface where heat is unphysically manufactured or destroyed.

599 The third inconsistent aspect is a direct consequence of the second; namely that if 600 one is tempted to post-calculate Conservative Temperature Θ from the model's 601 (assumed) values of potential temperature, the rate of change of the calculated ocean 602 heat content as the volume integral of $\rho c_n^0 \Theta$ would no longer be accurately related to the

603 heat that the atmosphere exchanged with the ocean. Neither would the area integral 604 between latitude bands of the air-sea heat flux be exactly equal to the difference between 605 the calculated oceanic meridional heat transports that cross those latitudes. Rather, 606 during the running of the model the heat that was lost from the atmosphere actually 607 shows up in the ocean as the volume integral of the model's prognostic temperature 608 variable. Thus we agree with Appendix D3.3 of Griffies et al. (2016) and strongly 609 recommend that Conservative Temperature is not calculated a posteriori in order to 610 evaluate heat content and heat fluxes in these EOS-80 based models.

611

612 4.1.2 Quantifying the air-sea flux imbalance

613 Here we quantify the air-sea flux errors involved with assuming that $T_{\rm model}$ of 614 EOS-80 models is potential temperature. These EOS-80 based models calculate the air-615 sea flux of their model's temperature as the air-sea heat flux, Q, divided by c_p^0 . However, since the isobaric specific heat capacity of seawater at 0 dbar is $c_p(S_*, \theta, 0)$, the 616 617 flux of potential temperature into the surface of the ocean should be Q divided by $c_p(S_*, \theta, 0)$. So, if the model's temperature variable is interpreted as being potential 618 619 temperature, the EOS-80 model has a flux of potential temperature entering the ocean that is too large by the difference between these fluxes, namely by Q/c_n^0 minus 620 $Q/c_n(S_*,\theta,0)$. This means that the ocean has received a different amount of heat than the 621 atmosphere actually delivers to the ocean, with the difference, ΔQ , being $c_n(S_*, \theta, 0)$ 622 623 times the difference in the surface fluxes of potential temperature, namely (for the last 624 part of this equation, see Eqn. (A.12.3a) of IOC et al., 2010)

625
$$\Delta Q = Q \left(\frac{c_p(S_*, \theta, 0)}{c_p^0} - 1 \right) = Q \left(\tilde{\Theta}_{\theta} - 1 \right).$$
(6)

We plot this quantity from the pre-industrial control run of ACCESS-CM2 in Figure 5c and show it as a cell area-weighted histogram in Figure 5e (note that while these plots apply to EOS-80 based ocean models, to generate these plots we have actually used data from ACCESS-CM2 which is a mostly TEOS-10 compliant model). The calculation takes into account the penetration of shortwave radiation into the ocean but is performed using monthly averages of the thermodynamics quantities. The temperatures and salinities at which the radiative flux divergences occur are taken into account in this calculation. However, the result is little changed if the sea surface temperatures and salinities are used with the radiative flux divergence assumed to take place at the sea surface. Results from similar calculations performed using monthly and daily averaged quantities in ACCESS-OM2 (Kiss et al. 2020) ocean-only model simulations were similar, suggesting that correlations between sub-monthly variations are not significant in such a relatively coarse-resolution model.

639 ΔQ has an area-weighted mean value of 16 mWm⁻² and we know that this 640 represents the net surface flux of potential temperature required to balance the volume 641 integrated non-conservation of potential temperature in the ocean's interior (Tailleux (2015)). To put this value in context, 16 mW m^{-2} corresponds to 5% of the observed trend 642 643 of 300 mWm^{-2} in the global ocean heat content from 1955-2017 (Zanna et al. 2019). In 644 addition to this mean value of ΔQ_{i} , we see from Figure 5c that there are small regions 645 such as the equatorial Pacific and the western north Pacific where ΔQ is as large as the 646 area-averaged heat flux, 300 mW m⁻², that the ocean has received since 1955. These local 647 anomalies of air-sea flux, if they existed, would drive local variations in temperature. 648 However, these ΔQ values do not represent real heat fluxes. Rather they represent the 649 error in the air-sea heat flux that we make if we insist that the temperature variable in an 650 EOS-80 based ocean model is potential temperature, with the ocean receiving a surface 651 heat flux that is larger by ΔQ than the atmosphere delivers to the ocean. Figure 6 shows 652 the zonal integration of ΔQ_{ℓ} in units of W per degree of latitude.

653 Figure 5e shows that, with T_{model} being interpreted as potential temperature, 5% 654 of the surface area of the ocean needs a surface heat flux that is more than 135 mW m^{-2} 655 different to what the atmosphere gives to/from the ocean. This regional variation of ΔQ of approximately $\pm 100 \text{ mW m}^{-2}$ is consistent with the regional variations in the air-sea 656 657 flux of potential temperature found by Graham and McDougall (2013) that is needed to 658 balance the depth-integrated non-conservation of potential temperature as a function of latitude and longitude. Figures 5d,f show that much of this spread is due to the 659 660 variation of the isobaric specific heat capacity on salinity, with the remainder due to the 661 variation of this heat capacity with temperature. We note that if this analysis were

662 performed with a model that resolved individual rain showers and the associated 663 freshwater lenses on the ocean surface, then these episodes of very fresh water at the sea 664 surface would be expected to increase the calculated values of ΔQ . Interestingly, by 665 way of contrast, it is the variation of the isobaric heat capacity with temperature that 666 dominates (by a factor of four) the contribution of this heat capacity variation to the *area-*667 *mean* of ΔQ (with the contribution of salinity, ΔQ_s , in Figure. 5d, leading to an area 668 mean of 4 mWm⁻²), as originally found by Tailleux (2015).

669 While a heat flux error of $\pm 100 \text{ mW m}^{-2}$ is not large, it also not trivially small, and 670 it seems advisable to respect these fundamental thermodynamic aspects of the coupled 671 Earth system. We will see that this $\pm 100 \text{ mW m}^{-2}$ issue is simply avoided by realizing 672 that the temperature variable in these EOS-80 models is not potential temperature.

673 In Appendix A we enquire whether the way that EOS-80 models treat their fluid 674 might be made to be thermodynamically correct for a fluid other than seawater. We find 675 that it is possible to construct such a thermodynamic definition of a fluid with the aim 676 that its treatment in EOS-80 models is consistent with the laws of thermodynamics. This 677 fluid has the same specific volume as seawater for given values of salinity, potential 678 temperature and pressure, but it has different expressions for both enthalpy and 679 entropy. This fluid also has a different adiabatic lapse rate and therefore a different 680 relationship between in situ and potential temperatures. However, this exercise in 681 thermodynamic abstraction does not alter the fact that, as a model of the real ocean, and 682 with the temperature variable being interpreted as being potential temperature, the 683 EOS-80 models have ΔQ more heat arriving in the ocean than leaves the atmosphere.

684 Since CMIP6 is centrally concerned with how the planet warms, it is advisable to 685 adopt a framework where heat fluxes and their consequences are respected. That is, we 686 regard it as imperative to avoid non-conservative sources of heat at the sea surface. It is 687 the insistence that the temperature variable in EOS-80 based models is potential 688 temperature that implies that the ocean receives a heat flux from the atmosphere that is 689 larger by ΔQ than what the atmosphere actually exchanges with the ocean. Since there 690 are some areas of the ocean surface where ΔQ is as large as the mean rate of global 691 warming, Option 1 is not supportable. This situation motivates Option 2 where we

- 692 change the interpretation of the model's temperature variable from being potential693 temperature to Conservative Temperature even when using EOS-80.
- 694

695 **4.2 Option 2: interpreting the EOS-80 model's temperature as being Conservative**

696 Temperature

697 Under this option the ocean model's temperature variable is taken to be Conservative 698 Temperature Θ . The air-sea flux of potential enthalpy is then correctly ingested into the 699 ocean model using the fixed specific heat c_n^0 , and the mixing processes in the model 700 correctly conserve Conservative Temperature. Hence the second, fourth and fifth items 701 listed in section 2 are handled correctly, except for the following caveat. In the coupled 702 model, the bulk formulae that set the air-sea heat flux at each time step use the 703 uppermost model temperature as the sea surface temperature as input. So with the 704 Option 2 interpretation of the model's temperature variable as being Conservative 705 Temperature, these bulk formulae are not being fed the SST (which at the sea surface is 706 equal to the potential temperature θ). The difference between these temperatures is 707 $\Theta - \theta$, which is the negative of what we plot in Figure 2. This is a caveat with this 708 Option 2 interpretation, namely that the bulk formula that the model uses to determine 709 the air-sea flux at each time step is a little different to what was intended when the parameters of the bulk formulae were chosen. This is a caveat regarding what was 710 711 intended by the coupled modeler, rather than what the coupled model experienced. 712 That is, with this Option 2 interpretation, the air-sea heat flux, while being a little bit 713 different than what might have been intended, does arrive in the ocean properly; there is 714 no non-conservative production or destruction of heat at the air-sea boundary as there is 715 in Option 1.

Regarding the remaining two items involving temperature listed in section 2, we can dismiss the fifth item, since any small difference in the initial values, set at the beginning of the lengthy spin-up period, between potential temperature and Conservative Temperature will be irrelevant after the long spin-up integration.

This then leaves the first point, namely that the model used the equation of state that expects potential temperature as its temperature input, $\tilde{\rho}(S_*/u_{\rm PS},\theta,p)$, but under this Option 2 we are interpreting the model's temperature variable as being Conservative Temperature. In the remainder of this section we address the magnitude of this effect, namely, the use of $\tilde{\rho}(S_*/u_{\rm PS},\Theta,p)$ versus the correct density $\tilde{\rho}(S_*/u_{\rm PS},\theta,p)$ which is almost the same as $\hat{\rho}(S_*,\Theta,p)$. Note, as discussed in section 3 above, the salinity argument of the TEOS-10 equation of state is taken to be S_* while that of the EOS-80 equation of state is taken to be $S_*/u_{\rm PS}$. These salinity variables are simply proportional to each other, and they have the same influence in both equations of state.

729 Under this Option 2 we are interpreting the model's temperature variable as 730 being Conservative Temperature, and so the density value that the model calculates 731 from its equation of state is deemed to be $\tilde{\rho}(S_*/u_{\rm PS},\Theta,p)$ whereas the density should be 732 evaluated as $\hat{\rho}(S_*, \Theta, p)$ where we remind ourselves that the hat over the *in situ* density 733 function indicates that this is the TEOS-10 equation of state, written with Conservative 734 Temperature as its temperature input. To be clear, under EOS-80 and under TEOS-10 735 the in situ density of seawater of Reference Composition has been expressed by two 736 different expressions,

751

$$\rho = \tilde{\rho} \left(S_* / u_{\rm PS}, \theta, p \right) = \hat{\rho} \left(S_*, \Theta, p \right), \tag{7}$$

both of which are very good fits to the *in situ* density (hence the equals signs); the increased accuracy of the TEOS-10 equation for density was mostly due to the refinement of the salinity variable, and the increase in the accuracy of TEOS-10 versus EOS-80 for Standard Seawater (Millero et al., 2008) was minor by comparison except for brackish seawater.

We need to ask what error will arise from calculating *in situ* density in the model as $\tilde{\rho}(S_*/u_{\rm PS},\Theta,p)$ instead of as the correct TEOS-10 version of in situ density, $\hat{\rho}(S_*,\Theta,p)$? The effect of this difference on calculations of the buoyancy frequency and even the neutral tangent plane is likely small, so we concentrate on the effect of this difference on the isobaric gradient of *in situ* density (the thermal wind).

Given that under this Option 2 the model's temperature variable is being
interpreted as Conservative Temperature, Θ, the model-calculated isobaric gradient of *in situ* density is

$$\tilde{\rho}_{S_*} \nabla_P S_* + \tilde{\rho}_{\theta} \nabla_P \Theta, \qquad (8)$$

752 whereas the correct isobaric gradient of *in situ* density is actually

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$$\hat{\rho}_{S_*} \nabla_P S_* + \hat{\rho}_{\Theta} \nabla_P \Theta. \tag{9}$$

Notice that here and henceforth we drop the scaling factor u_{PS} from the gradient expressions such as Eqn. (8). In any case, this scaling factor cancels from the expression, but we simply drop it for ease of looking at the equations; we can imagine that the EOS-80 equation of state is written in terms of S_* (which would simply require that a first line is added to the computer code which divides the salinity variable by u_{PS}).

The model's error in evaluating the isobaric gradient of *in situ* density is then thedifference between the two equations above, namely

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$$\operatorname{error} \operatorname{in} \nabla_{P} \rho = \left(\tilde{\rho}_{S_{*}} - \hat{\rho}_{S_{*}}\right) \nabla_{P} S_{*} + \left(\tilde{\rho}_{\theta} - \hat{\rho}_{\Theta}\right) \nabla_{P} \Theta .$$
(10)

The relative error here in the temperature derivative of the equations of state can bewritten approximately as

764
$$(\tilde{\rho}_{\theta} - \hat{\rho}_{\Theta})/\hat{\rho}_{\Theta} = \tilde{\alpha}^{\theta}/\hat{\alpha}^{\Theta} - 1,$$
 (11)

which is the difference from unity of the ratio of the thermal expansion coefficient with respect to potential temperature to that with respect to Conservative Temperature. This ratio, $\tilde{\alpha}^{\theta}/\hat{\alpha}^{\Theta}$, can be shown to be equal to $c_p(S_*,\theta,0)/c_p^0$ and we know (from Figure 1) that this varies by 6% in the ocean. This ratio is plotted in Figure 7(a). In regions of the ocean that are very fresh, a relative error in the contribution of the isobaric temperature gradient to the thermal wind will be as large as 6% while in most of the ocean this relative error will be less than 0.5%.

Now we turn our attention to the relative error in the salinity derivative of theequation of state, which, from Eqn. (10) can be written approximately as

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$$\left(\tilde{\rho}_{S_*} - \hat{\rho}_{S_*}\right) / \hat{\rho}_{S_*} = \tilde{\beta}^{\theta} / \hat{\beta}^{\Theta} - 1, \qquad (12)$$

and the ratio, $\tilde{\beta}^{\theta}/\hat{\beta}^{\Theta}$, has been plotted (at p = 0 dbar) in Figure 7(b). This figure shows that the relative error in the salinity derivative, $(\tilde{\rho}_{S_*} - \hat{\rho}_{S_*})/\hat{\rho}_{S_*}$, is an increasing (approximately quadratic) function of temperature, being approximately zero at 0°C, 1% error at 20°C and 2% error at 30°C. An alternative derivation of these implications of Eqn. (10) is given in Appendix B. 780 We conclude that under Option 2, where the temperature variable of an EOS-80 781 based model (whose polynomial equation of state expects to have potential temperature 782 as its input temperature) is interpreted as being Conservative Temperature, there are 783 persistent errors in the contribution of the isobaric salinity gradient to the isobaric 784 density gradient that are approximately proportional to temperature squared, with the 785 error being approximately 1% at a temperature of 20°C (mostly due to the salinity 786 derivative of *in situ* density at constant potential temperature being 1% different to the 787 corresponding salinity derivative at constant Conservative Temperature). Larger 788 fractional errors in the contribution of the isobaric temperature gradient to the thermal 789 wind equation do occur (of up to 6%) but these are restricted to the rather small volume 790 of the ocean that is quite fresh.

791 In Figure 8 we have evaluated how much the meridional isobaric density 792 gradient changes in the upper 1000 dbar of the World Ocean when the temperature 793 argument in the expression for density is switched from θ to Θ . As explained above, 794 this switch is almost equivalent to the density difference between calling the EOS-80 and 795 the TEOS-10 equations of state, using the same numeric inputs for each. We find that 796 19% of this data has the isobaric density gradient changed by more than 1% when 797 switching from θ to Θ . The median value of the percentage error is 0.22%; that is, 50% 798 of the data shallower than 1000 dbar has the isobaric density gradient changed by more 799 than 0.22% when switching from using EOS-80 to TEOS-10, with the same numerical 800 temperature input, which we are interpreting as being Θ .

Figure 8 should not be interpreted as being the extra error involved with taking T_{model} to be Conservative Temperature in EOS-80 ocean models, because, due to the lack of interior non-conservative source terms, the interpretation of T_{model} as being potential temperature is already incorrect by an amount that scales as Θ minus θ . Rather, Figure 8 illustrates the error in an EOS-80 model due to the use of an equation of state that is not appropriate to the way that its temperature variable is treated in the model.

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810 **4.3** Evaluating the options for EOS-80 models

811 Under option 1 where T_{model} is interpreted as potential temperature, there is a 812 non-conservation of heat at the sea surface, with the ocean seeing one heat flux, and the 813 atmosphere immediately above it seeing another, with 5% of the differences in these 814 heat fluxes being larger than approximately $\pm 100 \text{ mW m}^{-2}$, with a net imbalance of 815 16 mW m^{-2} .

816 Under option 2 where T_{model} is interpreted as Conservative Temperature, the air-817 sea flux imbalance does not arise, but two other inaccuracies arise. First, under option 2 818 the bulk formulae that determine part of the air-sea flux is based on the surface values of 819 Θ rather than of θ (for which the bulk formulae are designed). Second, the isobaric 820 density gradient in the upper ocean is typically different by ~1% to the isobaric density 821 gradient that would be found if the TEOS-10 equation of state had been adopted in these 822 models. These two aspects of option 2 are considered less serious than not conserving heat at the sea surface by up to $\pm 100 \text{ mWm}^{-2}$. Neither of the two inaccuracies that arise 823 824 under option 2 are fundamental thermodynamic errors. Rather they are equivalent to 825 the ocean modeler choosing (i) a slightly different bulk formulae, and (ii) a slightly 826 different equation of state. The constants in the bulk formulae are very poorly known so 827 that the switching from θ to Θ in their use will be well within their uncertainty (Cronin 828 et al., 2019) while the ~1% change to the isobaric density gradient due to using the 829 different equations of state is at the level of our knowledge of the equation of state of sea 830 water (see the discussion section below).

831 We conclude that option 2 where the T_{model} in EOS-80 models is interpreted as 832 Conservative Temperature is much preferred as it treats the air-sea heat flux in a manner 833 consistent with the First Law of Thermodynamics, and the treatment of T_{model} as being a 834 conservative variable in the ocean interior is more consistent with it being Conservative 835 Temperature than being potential temperature. These same two features of ocean models mean that T_{model} cannot be accurately interpreted as potential temperature, since 836 837 both the surface flux boundary condition and the lack of the non-conservative source terms in the ocean interior mean that these ocean models continually force $T_{\rm model}$ away 838 839 from being potential temperature, even if it was initialized as such.

841 **5.** Comparison with ocean observations

Now that we have argued that T_{model} of EOS-80 based models should be interpreted as being Conservative Temperature, how then should the model-based estimates of ocean heat content and ocean heat flux be compared with ocean observations and ocean atlas data? The answer is by evaluating the ocean heat content correctly in the observed data sets using TEOS-10, whereby the observed data is used to calculate Conservative Temperature, and this is used together with c_p^0 to evaluate ocean heat content and meridional heat fluxes.

849 We have made the case that the salinity variable in CMIP ocean models that have 850 been spun up for several centuries is Preformed Salinity S_{*} for the TEOS-10 compliant 851 models, and is $S_*/u_{\rm PS}$ for the EOS-80 compliant models. Hence it is the value of either 852 S_* or $S_*/u_{\rm PS}$ calculated from ocean observations to which the model salinities should be 853 compared. Preformed Salinity S_* is different to Reference Salinity S_R by only the ratio 854 0.26 = 0.35/1.35 compared with the difference between Absolute Salinity and Preformed 855 Salinity (see Figure 4), and these differences are generally only significantly different to 856 zero at depths exceeding 500 m. Note that Preformed Salinity can be evaluated from observations of Practical Salinity using the Gibbs SeaWater (GSW) software 857 858 gsw_Sstar_from_SP.

859

860 6. Discussion and Recommendations

861 We have made the case that it is advisable to avoid non-conservative sources of 862 heat at the sea surface. It is the prior interpretation of the temperature variable in EOS-863 80 based models as being potential temperature that implies that the ocean receives a 864 heat flux that is larger by ΔQ than the heat that is lost from the atmosphere. Since there 865 are some areas of the ocean surface where ΔQ is as large as the mean rate of global 866 warming, the issue is important in practice. This realization has motivated the new 867 interpretation of the prognostic temperature of EOS-80 ocean models as being 868 Conservative Temperature (our option 2, section 4.2).

869 A consequence of this new interpretation of the prognostic temperature variable 870 of all CMIP ocean models as being Conservative Temperature means that the EOS-80 871 based models suffer a relative error of ~1% in their isobaric gradient of *in situ* density in 872 the warm upper ocean. How worried should we be about this error? One perspective 873 on this question is to simply note (from above) that there are larger relative errors 874 (~2.7%) in the thermal wind equation in the deep ocean due to the neglect of variations 875 in the relative composition of sea salt. Another perspective is to ask how well science 876 even knows the thermal expansion coefficient, for example. From appendices K and O 877 of IOC et al. (2010) (and section 7 of McDougall and Barker (2011)) we see that the RMS 878 value of the differences between the individual laboratory-based data points of the 879 thermal expansion coefficient and the thermal expansion coefficient obtained from the fitted TEOS-10 Gibbs function is $0.73x10^{-6}$ K⁻¹ which is approximately 0.5% of a typical 880 881 value of the thermal expansion coefficient in the ocean. Without a proper estimation of 882 the number of degrees of freedom represented by the fitted data points, we might 883 estimate the relative error of the thermal expansion coefficient obtained from the fitted 884 TEOS-10 Gibbs function as being half of this, namely 0.25%. So a typical relative error in 885 the isobaric density gradient of ~1% in the upper ocean due to using Θ rather than θ as 886 the temperature input seems undesirable but not serious.

887 We must also acknowledge that all models have ignored the difference between 888 Preformed Salinity, Reference Salinity and Absolute Salinity (which is the salinity 889 variable from which density is accurately calculated). As discussed in IOC et al. (2010), 890 Wright et al. (2011) and McDougall and Barker (2011), glossing over these issues of the 891 spatially variable composition of sea salt, which is the same as glossing over the effects 892 of biogeochemistry on salinity and density, means that all our ocean and climate models 893 have errors in their thermal wind (vertical shear of horizontal velocity) that globally 894 exceed 2.7% for half the ocean volume deeper than 1000 m. In the deep North Pacific 895 Ocean, the misestimation of thermal wind is many times this 2.7% value. The 896 recommended way of incorporating the spatially varying composition of seawater into 897 ocean models appears as section A.20 in the TEOS-10 Manual (IOC et al. (2010), and as 898 section 9 in the McDougall and Barker (2012), with ocean models needing to carry a

899 second salinity type variable. While it is true that this procedure has the effect of 900 relaxing the model towards the non-standard seawater composition of today's ocean, it 901 is clearly advantageous to make a start with this issue by incorporating the non-902 conservative source terms that apply to the present ocean rather than to continue to 903 ignore the issue altogether. As explained in these references, once the modelling of 904 ocean biogeochemistry matures, the difference between the various types of salinity can 905 be calculated in real time in an ocean model without the need of referring to historical 906 data.

907 Nevertheless, we acknowledge that no published ocean model to date has 908 attempted to include the influence of biogeochemistry on salinity and density, and 909 therefore we recommend that the salinity from both observations and model output be 910 treated as Preformed Salinity S_* .

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2 6.1 Contrasts to the recommendations of Griffies et al. (2016)

913 How does this paper differ from the recommendations in Griffies et al. (2016)? 914 That paper recommended that the ocean heat content and meridional transport of heat 915 should be calculated using the model's temperature variable and the model's value of c_{p}^{0} , and we strenuously agree. However, in the present paper we argue that the 916 917 temperature variable carried by an EOS-80 based ocean model should be interpreted as 918 being Conservative Temperature, and not be interpreted as being potential temperature. 919 This idea was raised as a possibility in Griffies et al. (2016), but the issue was left unclear 920 in that paper. For example, section D2 of Griffies et al. (2016) recommends that TEOS-10 921 based models archive potential temperature (as well as their model variable, 922 Conservative Temperature) "in order to allow meaningful comparisons" with the output 923 of the EOS-80 based models. We now disagree with this suggestion; the thesis of the 924 present paper is that the temperature variables of both EOS-80 and TEOS-10 based 925 models are already directly comparable, and they should both be interpreted as being 926 Conservative Temperature, and they should both be compared with Conservative 927 Temperature from observations. The fact that the model's temperature variable is 928 labelled "thetao" in EOS-80 models and "bigthetao" in TEOS-10 based models we now

929 see as very likely to cause confusion, since we are recommending that the temperature
930 outputs of both types of ocean models should be interpreted as Conservative
931 Temperature.

932 The present paper also diverges from Griffies et al. (2016) in the way that the 933 salinity variables in CMIP ocean models should be interpreted and thus compared to 934 observations. Griffies et al. (2016) interpret the salinity variable in TEOS-10 based ocean 935 models as being Reference Salinity S_{R} whereas we show that these models actually 936 carry Preformed Salinity S_* but have errors in their calculation of densities. Similarly, 937 Griffies et al. (2016) interpret the salinity variable in EOS-80 based ocean models as being 938 Practical Salinity $S_{\rm p}$ whereas we show that these models actually carry $S_*/u_{\rm PS}$, that is, 939 Preformed Salinity divided by the constant, u_{PS} . This distinction between the present 940 paper and Griffies et al. (2016) is negligible in the upper ocean where Preformed Salinity 941 is almost identical to Reference Salinity (because the composition of seawater in the 942 upper ocean is close to Reference Composition), but in the deeper parts of the ocean, the 943 distinction is not negligible; for example, based on the work of McCarthy et al. (2015) we 944 have shown that the use of Absolute Salinity versus Preformed Salinity leads to ~1 Sv 945 difference in the meridional overturning streamfunction in the North Atlantic at a depth 946 of 2700 m. However, in this deeper part of the ocean, even though the difference 947 between Absolute Salinity and Preformed Salinity is not negligible, the difference 948 between Preformed Salinity and Reference Salinity (which the TEOS-10 based ocean 949 models have to date assumed their salinity variable to be) is smaller in the ratio 0.35/1.35 950 = 0.26 (see Figure 4). That is, if the salinity output of a TEOS-10 based ocean model was 951 taken to be Reference Salinity, the error would be only a quarter of the difference 952 between Absolute Salinity and Preformed Salinity, a difference which limits the 953 accuracy of the isobaric density gradients in the deeper parts of ocean models (see 954 Figure 4). A similar remark applies to EOS-80 based ocean models if their salinity 955 output is regarded as being Practical Salinity instead of being (as we propose) $S_*/u_{\rm PS}$.

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959 6.2 Summary table of ocean heat content imbalances

960 In Table 1 we summarize the effects of uncertainties in physical or numerical processes 961 in estimating ocean heat content or its changes. The first two rows are the rate of warming (expressed in mWm⁻² averaged over the sea surface) due to anthropogenic 962 963 global warming, and due to geothermal heating. The third row is an estimate of the 964 surface heat flux equivalent of the depth-integrated rate of dissipation of turbulent 965 kinetic energy, and the fourth is an estimate of the neglected net flux of potential 966 enthalpy at the sea surface due to the evaporation and precipitation of water occurring 967 at different temperatures.

968 The next (fifth) row is the consequence of considering the scenario where all the 969 radiant heat is absorbed into the ocean at a pressure of 25 dbar rather than at the sea 970 surface. The derivative of specific enthalpy with respect to Conservative Temperature at 25 dbar, \hat{h}_{Θ} , is c_p^0 times the ratio of the absolute in situ temperature at 25 dbar, $(T_0 + t)$, 971 to the absolute potential temperature, $(T_0 + \theta)$ at this pressure (see Eqn. (A.11.15) of IOC 972 et al. (2010)). The ratio of \hat{h}_{Θ} to c_p^0 at 25 dbar is typically different to unity by $6x10^{-6}$, 973 and taking a typical rate of radiative heating of 100 Wm⁻² over the ocean's surface leads 974 to 0.6 mWm⁻² as the area-averaged rate of mis-estimation of the surface flux of 975 976 Conservative Temperature for this assumed pressure of penetrative radiation. Since this is so small, the use of c_n^0 (rather than \hat{h}_{Θ}) to convert the divergence of the radiative heat 977 978 flux into a flux of Conservative Temperature is well supported, providing the correct 979 diagnostics are used for the calculation (such diagnostic issues may be responsible for 980 the heat budget closure issues identified by Irving et al. 2020).

The next six rows of Table 1 list the mean and twice the standard deviation of the volume integrated non-conservative production of Conservative Temperature, potential temperature, and specific entropy, all expressed in mWm⁻² at the sea surface. The following two rows are the results we have found in this paper for the air-sea heat flux error that is made if the EOS-80's temperature is taken to be potential temperature.

986 The final three rows show that ocean models, being cast in flux divergence form 987 with heat fluxes being passed between one grid box and the next, do not have appreciable numerical errors in deducing air-sea fluxes from changes in the volumeintegrated heat content.

990 The estimate from Graham and McDougall (2013) of -10 mWm^{-2} is for the net 991 interior production of θ , so this is a net destruction. A steady state requires this amount 992 of extra flux of θ at the sea surface (so it can be consumed in the interior). Our estimate 993 of this extra flux of θ at the sea surface is 16 mWm⁻², which is only a little larger than the 994 estimate of Graham and McDougall (2013).

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996 **6.3** Summary of recommendations

997 In summary, this paper has argued for the following guidelines for analyzing the998 CMIP model runs. We should

999 1. interpret the prognostic temperature variable of all CMIP models (whether they
1000 are based on the EOS-80 or the TEOS-10 equation of state) as being Conservative
1001 Temperature,

compare the model's prognostic temperature with the Conservative
 Temperature, Θ, of observational data,

1004 3. calculate the ocean heat content as the volume integral of the product of 1005 (i) in situ density (for non-Boussinesq models or reference density for 1006 Boussinesq) (ii) the model's prognostic temperature, Θ , and (iii) the model's 1007 value of c_n^0 ,

10084. interpret the salinity variable of the model output as being Preformed Salinity S_* 1009for TEOS-10 based ocean models, and S_*/u_{PS} for EOS-80 based ocean models (so1010it is advisable to post-multiply the salinity output of EOS-80 models by u_{PS} in1011order to have the salinity outputs of all types of CMIP models as Preformed1012Salinity S_*) and,

10135. compare the model's salinity variable with Preformed Salinity, S_* , calculated1014from ocean observations.

1015 6. Sea surface temperature should be taken as the model's prognostic temperature 1016 in the case of EOS-80 models (since this is the temperature that was used in the 1017 bulk formulae), and as the calculated and stored values of potential temperature1018 in the case of TEOS-10 models.

10197. Ensure that all required fixed variables, such as c_p^0 , (boussinesq) reference1020density, seawater volume, and freezing equation are saved to the CMIP archives1021alongside the prognostic temperature and salinity variables, so that analysts have1022all components required to accurately interpret the model fields. In addition,1023providing the full-depth OHC timeseries for each simulation would provide a1024quantified target for analysts to compare and contrast changes across models and1025simulations.

1026 Note that this sixth recommendation for EOS-80 based models exposes an unavoidable 1027 inconsistency in that the surface values of the model's prognostic temperature is best 1028 regarded internally in the ocean model as being Conservative Temperature, but we 1029 cannot avoid the fact that this same temperature was used as the sea surface (*in situ*) 1030 temperature in the bulk formulae during the running of such ocean models. Issues such 1031 as these will not arise when all ocean models have been converted to the TEOS-10 1032 equation of state.

1033 How then should the model's salinity and temperature outputs, S_* and Θ , be used to 1034 evaluate dynamical concepts such as streamfunctions, dynamic height, etc? The answer 1035 most consistent with the running of a numerical model is to use the equation of state 1036 that the model used, together with the model's temperature and salinity outputs on the 1037 native grid of the model. This method is important when studying detailed dynamical 1038 balances in ocean model output. But since we now have the output salinity and 1039 temperature of both EOS-80 and TEOS-10 models being the same (namely S_* and Θ), 1040 there is an efficiency and simplicity argument to analyze the output of all these models 1041 in the same manner, using algorithms from the Gibbs SeaWater (GSW) Oceanographic 1042 Toolbox of TEOS-10 (McDougall and Barker, 2011). Doing these model inter-1043 comparisons often involves interpolating the model outputs to different depths (or 1044 pressures) than those used in the original ocean model, so incurring some interpolation 1045 errors. While the use of the GSW software means that the in situ density will be 1046 calculated slightly differently than in some of the forward models, thus affecting the 1047 thermal wind and sea-level rise, these differences are small, as can be seen by comparing 1048 Figures A.5.1 and A.5.2 of the TEOS-10 Manual, IOC et al. (2010). Hence we think that it 1049 is viable for most purposes to evaluate density and dynamic height using the GSW 1050 Oceanographic Toolbox, with the input salinity to this GSW code being the model's 1051 Preformed Salinity, and the temperature input being the Conservative Temperature, 1052 which as we have argued, are the model's prognostic salinity and temperature variables. 1053 Another issue that may arise is where a TEOS-10 based model has been run with 1054 Conservative Temperature, but the monthly-mean Conservative Temperature output 1055 has been converted into potential temperature before sending the model output to the 1056 CMIP archive. What is the damage done if this inaccurately averaged value of potential 1057 temperature is converted back to Conservative Temperature using only the monthly-1058 mean potential temperature and salinity? While such an issue is perhaps an operational 1059 detail that takes us some distance from our intention of writing an academic paper about 1060 these issues, nevertheless we show Figure 9 which indicates that transforming between 1061 these monthly-averaged values is not a serious issue for relatively coarse-resolution 1062 ocean models.

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1064 Author Contribution

T J McD. devised this new way of interpreting CMIP ocean model variables, P. M. B and
R. M. H. provided figures for the paper, and all authors contributed to the concepts and
the writing of the manuscript.

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1078 Appendix A: A non-seawater thermodynamic interpretation of Option 1

1079 Ocean models have always assumed a constant isobaric heat capacity and have 1080 traditionally assumed that the model's temperature variable is whatever temperature 1081 the equation of state was designed to accept. Here we enquire whether there is a way of 1082 justifying Option 1 thermodynamically in the sense that Option 1 would be totally 1083 consistent with thermodynamic principles for a fluid that is different to real seawater.

That is, we pursue the idea that these EOS-80 based ocean models are not actually models of seawater but are models of a slightly different fluid. We require a fluid that is identical to seawater in some respects, such as that it has the same dissolved material (Millero et al., 2008) and the same issues around Absolute Salinity, Preformed Salinity and Practical Salinity, and the same in situ density as real seawater (at given values of Absolute Salinity, potential temperature and pressure). But we require that the expression for the enthalpy of this new fluid is different to that of real seawater.

1091 The difference that we envisage between real seawater and this new fluid is that, 1092 at zero pressure, the enthalpy of the new fluid is given exactly by the constant value c_p^0 1093 times potential temperature θ . That is, for the new fluid, potential enthalpy h^0 is 1094 simply $c_p^0 \theta$ (as it would be for an ideal gas), and the air-sea interaction for this new fluid 1095 would be exactly as it occurs in the EOS-80 based models. Moreover, conservation of 1096 potential temperature is justified for this new fluid, and the density and thermal wind 1097 would also be correctly evaluated in these EOS-80 based models.

1098 The enthalpy of this new fluid is then given by (since $h_p = v$)

1099
$$\breve{h}(S_{A},\theta,p) = c_{p}^{0}\theta + \int_{P_{0}}^{P} \tilde{v}(S_{A},\theta,p') dP',$$
(A1)

1100 while the entropy of this new fluid needs to obey the consistency relationship, 1101 $\ddot{\eta}_{\theta} = \ddot{h}_{\theta}(p=0)/(T_0+\theta)$, which reduces to

1102
$$\vec{\eta}_{\theta} = \frac{c_p^0}{\left(T_0 + \theta\right)},$$
 (A2)

1103 where $T_0 = 273.15$ K is the Celsius zero point. This consistency relationship is derived 1104 directly from the Fundamental Thermodynamic Relationship (see Table P.1 of IOC et al., 1105 2010). Integrating Eqn. (A2) with respect to potential temperature at constant salinity1106 leads to the following expression for entropy that our new fluid must obey,

1107
$$\tilde{\eta}(S_{A},\theta) = c_{p}^{0} \ln\left(1+\frac{\theta}{T_{0}}\right) + a\left(\frac{S_{A}}{S_{SO}}\right) \ln\left(\frac{S_{A}}{S_{SO}}\right).$$
(A3)

The variation here with salinity is taken from the TEOS-10 Gibbs-function-derived expression for specific entropy which contains the last term in Eqn. (A3) with the coefficient *a* being $a = -9.310292413479596 \text{ J kg}^{-1} \text{ K}^{-1}$ (this is the value of the coefficient derived from the g_{110} coefficient of the Gibbs function (appendix H of IOC *et al.* (2010)), allowing for our version of the normalization of salinity, (S_A/S_{SO})). This term was derived by Feistel (2008) to be theoretically correct at vanishingly small Absolute Salinities.

1115 With these definitions, Eqns. (A1) and (A3), of enthalpy and entropy of our new 1116 fluid, we have completely defined all the thermophysical properties of the fluid (see 1117 Appendix P of IOC et al., 2010 for a discussion). Many aspects of the fluid are different 1118 to seawater, including the adiabatic lapse rate (and hence the relationship between in 1119 situ and potential temperatures), since the adiabatic lapse rate is given by $\Gamma = \bar{h}_{\theta P} / \bar{\eta}_{\theta}$ 1120 and while the numerator is the same as for seawater (since $\tilde{h}_{\theta P} = \tilde{h}_{\theta P} = \tilde{v}_{\theta}$), the 1121 denominator, $\bar{\eta}_{\theta}$, which is now given by Eqn. (A2), can be up to 6% different to the 1122 corresponding function, $\tilde{\eta}_{\theta}$, appropriate to real seawater.

1123 We conclude that this is indeed a conceptual way of forcing the EOS-80 based 1124 models to be consistent with thermodynamic principles. That is, we have shown that these EOS-80 models are not models of seawater, but they do accurately model a 1125 1126 different fluid whose thermodynamic definition we have given in Eqns. (A1) and (A3). 1127 This new fluid interacts with the atmosphere in the way that EOS-80 models have 1128 assumed to date, the potential temperature of this new fluid is correctly mixed in the 1129 ocean in a conservative fashion, and the equation of state is written in terms of the 1130 model's temperature variable, namely potential temperature.

Hence we have constructed a fluid which is different thermodynamically to
seawater, but it does behave exactly as these EOS-80 models treat their model seawater.
That is, we have constructed a new fluid which, if seawater had these thermodynamic

characteristics, then the EOS-80 ocean models would have correct thermodynamics,
while being able to interpret the model's temperature variable as being potential
temperature.

But this does not change the fact that in order to make these EOS-80 models thermodynamically consistent in this way we have ignored the real variation at the sea surface of the isobaric specific heat capacity; a variation that we know can be as large as 6%.

1141 Hence we do not propose this non-seawater explanation as a useful 1142 rationalization of the behaviour of EOS-80 based ocean models. Rather, it seems less 1143 dramatic and more climatically relevant to adopt the simpler interpretation of Option 2. 1144 Under this option we accept that the model is modelling actual seawater, that the 1145 model's temperature variable is in fact Conservative Temperature, and that there are 1146 some errors in the equation of state of these EOS-80 models that amount to errors of the 1147 order of 1% in the thermal wind relation throughout much of the upper (warm) ocean. 1148 That is, so long as we interpret the temperature variable of these EOS-80 based models 1149 as Conservative Temperature, they are fine except that they have used an incorrect 1150 equation of state; they have used $\tilde{\rho}$ rather than $\hat{\rho}$. Apart from this "error" in the ocean 1151 code, Option 2 is a consistent interpretation of the ocean model thermodynamics and 1152 dynamics. In ocean models there are always questions of how to parameterize ocean 1153 mixing. To this uncertain aspect of ocean physics, under Option 2 we add the less than 1154 desirable expression that is used to evaluate density in EOS-80 based ocean models in 1155 CMIP

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1158 Appendix B: An alternative derivation of Eqn. (10)

1159 Eqn. (10) is an expression for the error in the isobaric density gradient when 1160 Conservative Temperature is used as the input temperature variable to the EOS-80 1161 equation of state (which expects its input temperature to be potential temperature). An 1162 alternative accurate expression to Eqn. (9) for the isobaric density gradient is

1163 $\tilde{\rho}_{S_*} \nabla_{\!_P} S_* + \tilde{\rho}_{\theta} \nabla_{\!_P} \theta , \qquad (B1)$

and subtracting this from the incorrect expression, Eqn. (8), gives the followingexpression for the model's error in evaluating the isobaric gradient of in situ density,

1166 error in
$$\nabla_{p} \rho = \tilde{\rho}_{\theta} \nabla_{p} (\Theta - \theta)$$
. (B2)

1167 An approximate fit to the temperature difference, $\Theta - \theta$, as displayed in Figure 2 is

1168
$$\left(\Theta - \theta\right) \approx 0.05 \Theta \left(1 - \frac{S_{\rm A}}{S_{\rm SO}}\right) - 1.75 \times 10^{-3} \Theta \left(1 - \frac{\Theta}{25^{\circ} \rm C}\right),$$
 (B3)

and using this approximate expression in the right-hand side of Eqn. (B2) gives

1170
$$\frac{\operatorname{error in} \nabla_{p} \rho}{\tilde{\rho}_{\theta}} \approx \left[0.05 \left(1 - \frac{S_{*}}{S_{SO}} \right) - 1.75 \times 10^{-3} \left(1 - \frac{\Theta}{12.5^{\circ} C} \right) \right] \nabla_{p} \Theta - \frac{0.05}{S_{SO}} \Theta \nabla_{p} S_{*}.$$
(B4)

1171 The first part of this expression that multiplies $\nabla_{\rho}\Theta$ corresponds to the proportional 1172 error in the thermal expansion coefficient displayed in Figure 7(a). The second part of 1173 Eqn. (B4) amounts to an error in the saline derivative of the equation of state, with the 1174 proportional error (corresponding to Eqn. (12)), being $-0.05 \tilde{\rho}_{\theta} \Theta / (\hat{\rho}_{S_A} S_{SO})$, and this is 1175 close to the error that can be seen in Figure 7(b). This error is approximately a quadratic 1176 function of temperature since the thermal expansion coefficient $\tilde{\rho}_{\theta}$ is approximately a 1177 linear function of temperature.

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	Heat flux contributions of different processes	mWm ⁻²
Physical processes	Global warming imbalance (Zanna et al., 2019), mean	+300
	Geothermal heating (Emile-Geay and Madec, 2009), mean	+86
	Viscous dissipation (Graham and McDougall, 2013), mean	+3
	Atmospheric water fluxes of enthalpy (Griffies et al. 2016), mean	- (150- 300)
Non-		
conservation errors	Extra flux of Θ if the air-sea radiative heat flux is taken to occur at a pressure of 25dbar	-0.6
	non-conservation of Θ (Graham & McDougall 2013), mean	+0.3
	non-conservation of Θ (Graham & McDougall 2013), 2*rms	+1
	non-conservation of θ (Graham & McDougall 2013), mean	-10
	non-conservation of θ (Graham & McDougall 2013), 2*rms	±120
	non-conservation of η (Graham & McDougall 2013), mean	+380
	non-conservation of η (Graham & McDougall 2013), 2*rms	+1200
	Interpreting EOS-80 T as θ (ACCESS-CM2 estimate), mean	+16
	Interpreting EOS-80 T as θ (ACCESS-CM2 estimate), 2*rms	±135
Numerical		
errors	ACCESS-OM2 single time-step	±10^(-7)
	ACCESS-OM2 diagnosed from OHC snapshots	±0.001
	ACCESS-CM2 diagnosed from OHC monthly- averages	±0.03

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- 1185**Table 1:** Summary of the impact of various processes and modelling errors on the global
- 1186 ocean heat budget and its imbalance. All numbers are in units of mWm^{-2} . Numerical errors
- are diagnosed from either ACCESS-OM2 (machine precision errors) or ACCESS-CM2
- 1188 (associated with not having access to OHC snapshots). Numbers from interior processes are
- 1189 converted to equivalent surface fluxes by depth integration. The sign convention here is that a
- 1190 positive heat flux is heat entering the ocean or warming the ocean by internal dissipation. The
- 1191 symbol η in this table stands for entropy.

1193 Code Availability

- 1194 This paper has not run any ocean or climate models, and so has not produced any
- 1195 such computer code. Processed data and code to produce the ACCESS-CM2 figures 5,
- 1196 6 and 9 is located at the github repository
- 1197 <u>https://github.com/rmholmes/ACCESS_CM2_SpecificHeat.</u>
- 1198
- 1199

1200 Data Availability

- 1201 This paper has not produced any model data. Processed data and code to produce the
- 1202 ACCESS-CM2 figures 5, 6 and 9 is located at the github repository
- 1203 https://github.com/rmholmes/ACCESS CM2 SpecificHeat.
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Absolute Salinity, S_A (g kg⁻¹) **Figure 1.** (a) Contours of isobaric specific heat capacity c_p of seawater (in J kg⁻¹ K⁻¹), at p = 0 dbar. (b) a zoomed-in version for a smaller range of Absolute Salinity. The dashed line is the freezing line at p = 0 dbar.



Figure 2. Contours (in °C) of the difference between potential temperature and

- 1318 Conservative Temperature, $\theta \Theta$.
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Contours of $\Theta - \hat{h}(S_A, \Theta, 1000 \, \text{dbar})/c_p^0$ on the Absolute Salinity – 1323 Figure 3. $\hat{h}(S_A,\Theta,1000 \text{ dbar})/c_p^0$ diagram. Enthalpy, $\hat{h}(S_A,\Theta,1000 \text{ dbar})$, is a conservative 1324 1325 quantity for turbulent mixing processes that occur at a pressure of 1000dbar. The 1326 mean value of the contoured quantity is approximately -2.44°C illustrating that 1327 enthalpy does not possess the "potential" property; that is, enthalpy increases 1328 during adiabatic and isohaline increases in pressure. The fact that the contoured quantity on this figure is not a linear function of S_A and $\hat{h}(S_A,\Theta,1000 \text{ dbar})$ 1329 1330 illustrates the (small) non-conservative nature of Conservative Temperature. The 1331 dots are data from the word ocean at 1000dbar.





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Figure 4. Number line of salinity, illustrating the differences between Preformed Salinity $\,S_{\rm *}$, Reference Salinity $\,S_{\rm R}$, and Absolute Salinity $\,S_{\rm A}\,$ for seawater whose composition differs from that of Standard Seawater which has Reference Composition. If a seawater sample has Reference Composition, then $\delta S_A = 0$ and S_* , $S_{\rm R}$ and $S_{\rm A}$ are all equal.



0.01

0 ∟ -0.4

-0.3

-0.2

-0.1

0

Cell area-weighted ΔQ_S (Wm⁻²)

0.1

0.2

0.3

0.4

1348 1349

0.01

0 ∟ -0.4

-0.3

-0.2

-0.1

0

Cell area-weighted $\Delta Q \; (Wm^{-2})$

0.1

0.2

0.3

0.4

1350 Figure 5. (a) The average value of the ratio of the isobaric specific heat of 1351 seawater and c_p^0 for data from the ACCESS-CM2 model's pre-industrial 1352 control simulation (600 years long). (b) The average surface heat flux Q (Wm⁻ 1353 ²) in this same ocean model. (c) The additional heat that the ocean 1354 receives/loses compared to the heat that the atmosphere loses/receives 1355 (assuming that an EOS-80 model's temperature variable is potential 1356 temperature), ΔQ (Wm⁻², Eqn. 6). (e) a histogram of ΔQ weighted by the 1357 area of each grid cell. (d) The contribution of salinity variations to the air-sea heat flux discrepancy, given by $\Delta Q_s = Q(S-\overline{S})(1/c_p^0)\partial c_p/\partial S$, where \overline{S} is the 1358 surface mean salinity and $\partial c_n / \partial S$ is the variation in the specific heat with 1359 1360 salinity at the surface mean salinity and potential temperature. (f) a 1361 histogram of ΔQ_s weighted by the area of each grid cell. Shown in red in panels e and f are the mean, 5^{th} and 95^{th} percentiles of the histogram (Wm⁻²). 1362 1363 Note that these calculations neglect correlations between surface properties 1364 and the surface heat flux at sub-monthly time scales. 1365





Figure 6. The ACCESS-CM2 zonally integrated ΔQ From Fig.5c, showing the imbalance in the air-sea heat flux in Watts per degree of latitude.







1376 **Figure 7.** (a) The ratio of the thermal expansion coefficients with respect to Conservative 1377 Temperature and potential temperature, $\tilde{\alpha}^{\theta}/\hat{\alpha}^{\Theta} = \tilde{\Theta}_{\theta}$. (b) The ratio of the saline 1378 contraction coefficients at constant potential temperature to that at constant Conservative 1379 Temperature, $\tilde{\beta}^{\theta}/\hat{\beta}^{\Theta} = 1 + (\hat{\alpha}^{\Theta}/\hat{\beta}^{\Theta})\hat{\theta}_{S_{\star}}/\hat{\theta}_{\Theta}$ at p = 0 dbar.

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- 1381





1383 Figure 8. The northward density gradient at constant pressure (the horizontal axis) for 1384 data in the global ocean atlas of Gouretski and Koltermann (2004) for p < 1000 dbar. The 1385 vertical axis is the magnitude of the difference between evaluating the density gradient 1386 using Θ versus θ as the temperature argument in the expression for density. This is 1387 virtually equivalent to the density difference between calling the EOS-80 and the TEOS-10 1388 equations of state, using the same numeric inputs for each. The 1% and 2% lines indicate 1389 where the isobaric density gradient is in error by 1% and 2%. 19% of the data shallower 1390 than 1000 dbar has the isobaric density gradient changed by more than 1% when 1391 switching between the equations of state. The median value of the percentage error in the 1392 isobaric density gradient is 0.22%. 1393 1394



Figure 9. The RMS error (K) in evaluating Conservative Temperature from the CMIP6
archived monthly-averaged values of potential temperature and salinity, compared with
averaging the instantaneous values of Conservative Temperature for a month at the (a)
surface and (b) the zonal mean. These quantities are calculated from 50 years of
temporally averaged output from the ACCESS-CM2 model's pre-industrial control
simulation. The errors are seen to be no larger than a few mK.