Interactive comment on “Improved routines to model the ocean carbonate system: mocsy 1.0” by J. C. Orr and J.-M Epitalon

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1 General comments

1.1 Appreciation of the manuscript

In this paper, J. C. Orr and J.-M. Epitalon describe a new Fortran 95 library to model the ocean carbonate system, MOCSY. MOCSY updates and extends the widely used codes from the Ocean Carbon Cycle Model Intercomparison Project, OCMIP (Orr et al., 2000). It extends the choice of stoichiometric constant parameterizations, to comply, as far as possible with the recommendations for best practices (Dickson et al., 2007). Deviations from best practices are clearly stated and entirely justified. MOCSY further-
more provides more complete speciation calculations than the OCMIP codes.

MOCSY also offers its users the possibility to take into account developments that got published after the best practices. These include the new boron-to-salinity ratio from Lee et al. (2010) and more recent parameterizations for the two dissociation constants of carbonic acid. It furthermore goes beyond the recommendations for best practices, by also providing the necessary formulations for pressure correction (Dickson et al. (2007) does not include these since its focus is on the analytical aspects of carbonate system measurements, generally made at atmospheric pressure).

Finally, the paper also presents a quantitative assessment of several approximations often made in models: (1) density is constant; (2) the approximation of in-situ temperature by potential temperature (the former being, strictly speaking, the temperature actually required for the chemistry calculations, while the latter is what models provide); (3) the equivalence between depths in meters and pressure in dbar; (4) nutrient contributions to alkalinity (i.e., from the silicic and phosphoric acid systems) are negligible.

The paper is generally well written. The history and motivations behind the development of MOCSY are informative. In a few instances, the text is not entirely precise – this can, however, be easily fixed. The selection of material to demonstrate the potential of the package is good. Figures are of good quality, but might be improved for a few details (coordinate axes extents).

The model description is somewhat short. While more or less all of the thermodynamical ingredients are described in full detail, close to nothing is provided regarding the numerical methods applied to actually perform the pH calculations. Above all, the little that is provided is contradictory. Geoscientific Model Development papers are expected to present a complete and detailed description of the models. A minimum of technical details about the methods and algorithms used should be given and limitations should also be discussed (see “Manuscript Types” in the “Submission” guidelines).
The presentation and discussion of the results contain, unfortunately, a few inaccuracies (see detailed comments below). These do, however, not impinge on the overall conclusions and can be easily corrected.

The “Code availability” section in the paper and the code distribution channel are exemplary. The code is distributed under the MIT license, and can be easily obtained. The included test case compiled and worked out of the box as expected for me.

I am confident that the authors will be able to address all of the shortcomings mentioned. Accordingly, I recommend to publish this paper after a revision of the text.

1.2 Discussion

1.2.1 Constant, variable, or no density correction at all?

Regarding issue (1) above, I expect that applying a constant or even no density correction at all should not make much of a difference in the calculation results! This may seem counterintuitive, but, it is sufficient to notice the following:

1. all of the equilibrium constants (except for $K_W$) carry the units of the proton concentration – for all other concentrations that appear in the equilibrium relationships, any unit conversion factors cancel out;

2. in the alkalinity-pH equation, again, only the ratios of the different total concentrations (total dissolved inorganic carbon, $C_T$, total borate, $B_T$, sulfate, $S_T$, etc. . . .) to total alkalinity, $A_T$, are relevant and, as long as all of these are expressed relative to the same reference framework (volume or mass), any density conversion factors will cancel out; the only term in the alkalinity-pH equation that might be affected by an imprecise density will be $A_W/A_T$, where the water self-ionization alkalinity, $A_W$, directly depends on the proton concentration units. That ratio is, however, only of the order of a per mil and an error of the order of a percent in this actual ratio should not make any significant difference in the calculated proton concentration.

Whatever the units of the total input concentrations, the speciation results will not be affected to any significant extent, since the speciation routine only determines, e.g., what fraction of $C_T$ is $CO_2^\pm$, $HCO_3^-$ or $CO_3^{2-}$. Only the calculated proton concentration will always come out in the units of the equilibrium constants (generally mol/kg-soln) and it would be best not to convert it to a volumetric reference at all.

Using a different density conversion factor during subsequent model-data comparison than the one that was possibly used for the speciation calculation should not make any significant difference. The model-data comparison must anyway be carried out consistently at the density of the data samples.

1.2.2 Depth-pressure conversion: how significant are the errors that we try to avoid?

Regarding issue (3) above, Lewis and Wallace (1998) already noted that depth expressed in meters and pressure expressed in dbar differ by only 3% at 10,000 dbar and less at lower pressures, which is well within the uncertainties of the pressure effects on the equilibrium constants. To my best knowledge, there has not been any progress in reducing the uncertainties of the pressure correction coefficients since the compilation of Millero (1995) and the reviewing efforts of Lewis and Wallace (1998): the currently used coefficients are more than 30 years old; some of them remain are merely expert guesses – the coefficient for the first dissociation constant of silicic acid is estimated from that of boric acid (Millero, 1995).

Accordingly one may ask whether the tiny improvement resulting from the more accu-
rate depth-to-pressure conversion is really worth the effort, since the resulting changes
do not have any measurable significance.

Furthermore, the common practical application of the pressure correction of the equi-
librium constants leaves, in my opinion, an unpleasant aftertaste of inconsistency: for
any chosen work pH scale, to adjust the value of an equilibrium constant calculated
at atmospheric pressure to any other given pressure, Lewis and Wallace (1998) rec-
ommend to first convert it to the seawater (SWS) or the NBS scales, then apply the
pressure correction, and finally convert the result back to the selected work pH scale
(using the pH scale conversion factor at the final pressure). Since NBS scales are
nowadays rarely used in seawater chemistry, the pressure correction is generally made
on the SWS scale. Results will nevertheless most probably be different depending on
whether the SWS or the NBS pathways are chosen. This protocol must be followed
for all equilibrium constants, except for the dissociation of hydrogen sulfate and fluoric
acid, which should be pressure-adjusted on the free scale. These are indeed required
for the pH scale conversions, and the common reference in pH scale conversions is
the free pH scale.

While I see the usefulness of a well-defined protocol for performing these pressure
adjustments, we are now in the somewhat surrealist situation where the best practices
recommend the use of equilibrium constants on the total pH scale, these constants
need to be temporarily converted to the SWS scale for pressure correction, and back to
the total scale, except for two of them, that need to stick to the free scale, although the
pressure correction coefficients are certainly not precise enough to distinguish between
the different scales. Why not carry out the pressure correction on the free scale and
gain at least in consistency?

1.2.3 Nutrient alkalinity: how significant are the differences between the results?

Although, once again, the assessment and the discussion of the importance of nutrient
related acid systems in the alkalinity composition is very interesting, I am not convinced
about the significance of the consequences of including or neglecting them. As far as
I can see, the quality of even our most advanced global model results is still far from
sufficient to make the observed differences relevant. Locally in the deep-sea, inter-
model differences in the calculated carbonate ion profiles remain far larger than the
quoted 1.6 \( \mu \)mol/kg. A polynomial pH-solver based upon practical alkalinity may easily
be an order of magnitude faster (even if safe-guarded) than a full solver required with
the complete representation of \( A_T \).

Finally, using observed present-day nutrient climatologies for studies of the past and
future evolution of the ocean carbon cycle, where production-remineralization-burial
patterns differ from the present ones, may possibly lead to even larger errors than
those that we are trying to address here.

2 Specific comments

In the following, when reference is made to \textsc{CO2SYS}, it should be understood that it is
the MATLAB version that is meant (van Heuven et al., 2011), the only version for which
that I was able to inspect the source code.

Page 2880, lines 18–19: this is rather cryptic. Does this mean that the errors listed
by Lewis and Wallace (1998) for Millero (1995) have been taken into account? If so, it
would be helpful to make the text more precise.

Page 2881, line 6: \( \text{PO}_4^{3-} \) is ambiguous (is it the species or the total dissolved phos-
... phate that is meant?). I guess it should read “total phosphate”.

**Page 2881, lines 14–16:** it was previously said (p. 2880, ll. 18–19), that the pressure-correction coefficients came from Lewis and Wallace (1998). Here we find other references. Are those the same coefficients? If so, the original reference should be cited; if not, how do they differ? Please clarify.

**Page 2881, lines 10–12:** The description of the method used to solve the alkalinity-pH equation is insufficient and does not appear to be correct:

1. Maier-Reimer (1993) uses the practical alkalinity approximation and specifies that a Newton iteration is used to solve the alkalinity-pH equation, expressed as a variant of the rational function form (see, e.g., Munhoven (2013) for the different forms of the equation);
2. Aumont and Bopp (2006) describe the PISCES model, which, although it derives from HAMOCC5 (Aumont et al., 2003), claims to use the OCMIP protocol (i.e., Newton + bi-section) for its carbonate chemistry, with practical alkalinity as an approximation to total alkalinity;
3. Inspection of the MOCSY code shows that the adopted method is actually not a Newton method, as the cited literature might suggest, but, in the classification of Munhoven (2013), a fixed-point iterative carbonate alkalinity correction (ICAC) method, combined with a pH = 8 initialization scheme.

It would be worth noting that among the six different solution algorithms studied by Munhoven (2013), icacfp, which is equivalent to the ICAC algorithm adopted here, was the second-least efficient one, two to three times slower than the best (if we only consider the SW1 and SW2 benchmarks that it passed and disregard the SW3 benchmark that it failed). Unless the cubic initialization scheme of Munhoven (2013) was used, it also failed the RTC1 stress-test in about 40–90% of the cases, and the RTC2 stress-test in 77–100% of the cases (Munhoven, 2013, suppl.). The safe-guarded algorithms presented by Munhoven (2013), on the other hand, passed both stress-tests in 100% of the cases, and were at worst 15% slower than the fastest methods in the SW1, SW2, and SW3 benchmarks.

As further shown by Munhoven (2013), ICAC methods present inherent convergence problems at high $A_T / C_T$ ratios and at low $C_T$ values. These convergence problems are unavoidable and can be alleviated only to a very limited extent. There is no way to efficiently safeguard them, as it is the underlying recurrence that becomes divergent.

ICAC methods are fine for typical present-day conditions, although comparatively slowly converging. It is, however, risky to rely on them for conditions that deviate from present-day. They are, e.g., not the best choice in 3D models that are intended to be used to assess the effects of alkalinization, a geoengineering technique to mitigate ocean acidification and that may locally lead to extremely high $A_T$ concentrations, or the impact of melt water pulses, which might locally lead to too dilute $C_T$ for ICAC methods.

Many of these aspects and possible alternatives are covered in detail by Munhoven (2013).

The model description needs to be corrected, completed and possible limitations discussed. It could also be better streamlined: e.g., in the current manuscript, the pressure-correction of the equilibrium constants is mentioned in four different places (p. 2880, ll. 18–19; p. 2881, ll. 14–15; p. 2882, ll. 13–15; p. 2884, ll. 2–5), and still, only insiders are likely understand the implications.

**Page 2881, lines 24–28:** There are a few more exceptions than these three. Similarly to $K_S$, $K_F$ must also be referenced on the free scale (and that is actually how it is implemented in the code). $K_S$ is also independent on pH scale.
Page 2882, line 17: please notice that, if the calcium content is calculated following Riley and Tongudai (1967), as stated, then the correct Ca-to-chlorinity ratio is 0.02128 and not 0.02127, as reported by Dickson et al. (2007, chap. 5, tab. 2). The 0.02127 value has been around in the literature for a while without an adequate reference: it can at least be traced back to Millero (1982), where the cited original references do not allow an unequivocal attribution. I speculate that it was obtained by averaging the 0.02126 from Culkin and Cox (1966) and the 0.02128 from Riley and Tongudai (1967).

The difference is definitely small, but if confusion can be avoided...

Pages 2884–2885, section 3.1: Where do the diagnosed differences in the species concentrations come from? For identical input, I would expect no differences at all (at least to within an order of magnitude of machine precision or so). The differences are not random, so there must be some intrinsic differences between the ways MOCSY and CO2SYS carry out these calculations.

The reported discrepancies in the Revelle factor are quite interesting: the largest differences arise where the nutrient contributions to alkalinity are the largest. This points out an important shortcoming in MOCSY’s Revelle factor calculation: the implemented analytical buffer factor formulation of Frankignoulle (1994) was derived for the practical alkalinity approximation. CO2SYS, on the other hand, uses a finite difference approach to calculate the Revelle factor, using results derived with the (most) complete alkalinity representation. Despite being based upon a numerical approximation only, the CO2SYS approach might be more reliable, since it is consistent with the adopted $A_T$ approximation.

Pages 2885–2886, section 3.2: this section is not totally accurate and partially contradictory.

1. The 4% increase of the total borate concentration, $B_T$, does not translate into a 4% increase of borate alkalinity, $A_B$, as could be easily verified with, e.g., MOCSY. There is some buffering. Indeed, if total alkalinity, $A_T$, stays constant, together with all the other total concentrations, pH will adjust (decrease), such as to decrease all of the other alkalinity contributions besides $A_B$. Calculations with SolveSAPHE 1.0.1 (Munhoven, 2013) indicate that that 4% $B_T$ increase leads to $A_B$ increase of close to 3% (using the practical alkalinity approximation for the sake of simplicity).

2. The $B_T$ increase does not affect $A_T$ as stated, nor does it affect $C_T$ (also an input variable). The conclusion that “the $C_T$ must also decline” (p. 2886, l. 12) does therefore not make sense. While it is correct that carbonate alkalinity $A_C$ must decline, as a result of the increased $A_B$, this decline is brought about by a pH decrease, not by a $C_T$ change. The buffer capacity is dependent on the actual levels of all of the total dissolved acid concentrations, and on pH (which of course depends on all the previous). With a different pH, a different buffer capacity is obtained. It is difficult to predict the evolution of the buffer capacity for this change a priori, as its analytical expression involves a complicated product of factors that present counteracting variations.

3. For the same reasons (i.e., $A_T$ and $C_T$ are input variables), and contrary to what is written at ll. 12–13, $A_T$ minus $C_T$ remains constant. Here, the commonly made approximation $[\text{CO}_2^{3-}] \approx A_T - C_T$ is terribly misleading. It would be far more accurate to use $[\text{CO}_2^{3-}] \approx A_C - C_T$, which is more compatible with what is observed here.

Page 2886, line 16 – page 2886, line 2: issues related to the use of constant density are probably not as important as outlined here (see general comments above).

Pages 2887–2888: how significant are the calculated differences? Are inter-model differences not far larger than this?
3 Technical corrections

Page 2880, line 17: strange sentence “[...] recommended by for best practices [...]” – please check.

Page 2897, Fig. 3: the horizontal axes on the top left and on all of the lower panels should be modified to better separate the different curves. Much space is currently wasted.

4 Code

4.1 Availability and distribution

Two ways to get the code are indicated, together with a link to a dedicated website of the code, where extensive information is available. The common user who does not have git installed can easily go to the github page and download the zip archive.¹

I have downloaded, compiled and executed the code on a Linux machine with the gfortran compiler. Everything worked out of the box.

¹Subversion (svn) aficionados may use github’s svn-git bridge service, allowing them to use the favorite version control system and still have access to the complete history of the package. See github’s help pages for more details about this facility.

4.2 Comments on the code

1. Users will appreciate the extensive comments and references in the source code.

2. make test_mocsy produces a libmocsy.a library, which is nevertheless not used (not even linked into the produced test_mocsy).

3. The code is a mixture of single and double precision operations. While this is perfectly valid, it leads, in my personal opinion and in my experience as a teacher and instructor in Fortran programming, to unnecessary numerical artifacts. Why not use a uniform real type throughout, the more since the flexible typing mechanism offered by Fortran 90 and later is already used?

4. In some of the subroutines one reads in the comments that they are “Needed because xyz is a function (cannot accept arrays)”. While this was to some extent true in FORTRAN 77 (arguments could well be arrays, but results not), it is definitely not correct in the Fortran 90 and later standards. Those do also allow array-valued functions. In MOCSY, converting the functions from scalar to array-valued might be interesting for performance reasons, since most (if not all) of the concerned function/subroutine pairs only perform straightforward calculations (no iterations) and might thus take advantage of the efficient whole-array operations.

5. The Dickson and Riley $K_F$ is converted from the free to the total pH scale, to be converted back to the free scale right away before being corrected for pressure. The two conversion steps could of course be skipped (and the Perez and Fraga $K_F$ directly converted to the free scale instead).
References


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