# Package AquaEnv: an Aquatic modelling Environment in R

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#### Abstract

**AquaEnv** is an integrated development toolbox for aquatic chemical model generation focused on (ocean) acidification and CO2 air-water exchange.

- It contains all elements necessary to model the pH, the related CO2 air-water exchange, as well as aquatic acid-base chemistry in general for an arbitrary marine, estuarine or freshwater system. Also chemical batches can be modelled.
- Next to the routines necessary to calculate desired information, AquaEnv also contains a suite of tools to visualize this information.
- Furthermore, **AquaEnv** can not only be used to build dynamic models of aquatic systems, but it can also serve as a simple desktop tool for the experimental aquatic chemist to generate and visualize all possible derived information from a set of measurements with one single easy to use R function.
- Additionally, the sensitivity of the system to variations in the input variables can be visualized.
- AquaEnv also contains a number of example "applications" that make use of the aquatic modelling toolbox that AquaEnv provides:
  - a theoretical titration simulator
  - and a routine to determine total alkalinity ([TA]), the total dissolved inorganic carbon concentration ([ $\sum$ CO2]), as well as additionally the electrode standard potential (E<sub>0</sub>) and the first dissociation constant of the carbonate system ( $K_{CO_2}^*$ )

*Keywords*: aquatic modelling, pH, pH scales, dissolved inorganic carbon, total alkalinity, total alkalinity curve fitting, theoretical titration, revelle factor, omega, solubility products, CO<sub>2</sub>, ocean acidification, estuaries, carbonate system, seawater, R.

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# 1 Introduction

**AquaEnv** is a toolbox for aquatic modelling that serves several purposes

- It provides functions to calculate the stoichiometric equilibrium constants (K\*) for key acid base systems in natural seawater, the Henry's constants (K<sub>0</sub>), as well as the solubility products (K<sub>sp</sub>) for calcite and aragonite. This functionality is provided via the functions K\_CO2, K\_HCO3, K\_BOH3, K\_W, K\_HSO4, K\_HF, K\_NH4, K\_H2S, K\_H3PO4, K\_H2PO4, K\_HPO4, K\_SiOH4, K\_SiOOH3, KO\_CO2, KO\_O2, Ksp\_aragonite, and Ksp\_calcite.
- It is designed to make its use as easy as possible: all the information that can be calculated from the set of parameters know of a system or sample can be obtained by one single function: aquaenv. This function returns a list of class aquaenv that contains next to the input parameters
  - the clorinity, the ionic strength,  $[\sum B(OH)_3]$ ,  $[\sum H_2SO_4]$ ,  $[\sum HF]$ ,  $[Cl^-]$ ,  $[Cl^-]$ ,  $[\sum Br]$ ,  $[Na^+]$ ,  $[Mg^{2+}]$ ,  $[Ca^{2+}]$ ,  $[K^+]$ ,  $[Sr^{2+}]$  calculated from salinity as given in DOE (1994) (Please note that if values for  $[\sum B(OH)_3]$ ,  $[\sum H_2SO_4]$ ,  $[\sum HF]$  are given as input parameters, these parameters are used and not the ones calculated from salinity.)
  - the hydrostatic pressure calculated from the given depth and the seawater density calculated from temperature and salinity as given by Millero and Poisson (1981)
  - a set of conversion factors to convert between different pH scales (Dickson 1984;
     Zeebe and Wolf-Gladrow 2001) and between mol/kg-H<sub>2</sub>O and mol/kg-solution (inferred from Roy, Roy, Vogel, PorterMoore, Pearson, Good, Millero, and Campbell (1993b) and DOE (1994))
  - the Henry's constants for  $CO_2$  (Weiss 1974) and for  $O_2$  (inferred from Weiss 1970) calculated from temperature and salinity as well as the associated saturation concentrations of  $CO_2$  and  $O_2$ .
  - the ion product of water (Millero 1995), the stoichiometric equilibrium constants of HSO<sub>4</sub><sup>-</sup> (Dickson 1990a), HF(Dickson and Riley 1979a), CO<sub>2</sub> (Roy et al. 1993b), HCO<sub>3</sub><sup>-</sup> (Roy et al. 1993b), B(OH)<sub>3</sub> (Dickson 1990a), NH4<sup>+</sup>(Millero, Yao, and Aicher 1995), H2<sub>S</sub> (Millero 1995), H<sub>3</sub>PO4(Millero 1995), H<sub>2</sub>PO4<sup>-</sup> (Millero 1995), HPO4<sup>2-</sup> (Millero 1995), SiOH4 (Millero, Plese, and Fernandez 1988), SiOOH3<sup>-</sup> (Wischmeyer, Del Amo, Brzezinski, and Wolf-Gladrow 2003), HNO2 (Riordan, Minogue, Healy, O'Driscoll, and Sodeau 2005), HNO3, H2SO4 (Atkins 1996), HS (Atkins 1996) mostly calculated as functions of temperature and salinity and pressure corrected according to Millero (1995).
  - the solubility products of calcite and aragonite (Mucci 1983) as well as the associated  $\Omega$ 's if a full speciation is calculated (see below)
  - the partial pressure of CO<sub>2</sub> if a full speciation is calculated (see below)
  - if  $[\sum CO_2]$  and pH are given [TA] is calculated, if  $[\sum CO_2]$  and [TA] are given pH is calculated, if  $[\sum CO_2]$  and  $[CO_2]$  or pCO<sub>2</sub> are given, pH and [TA] are calculated.
  - if either one of the pairs pH and  $[CO_2]$  or pCO<sub>2</sub>, pH and [TA], or [TA] and  $[CO_2]$  or pCO<sub>2</sub> is given,  $[\sum CO_2]$  is calculated

- if sufficient information is given and the flag speciation=TRUE is set, a full speciation of  $[\sum CO2]$ ,  $[\sum NH4]$ ,  $[\sum H_2S]$ ,  $[\sum HNO3]$ ,  $[\sum HNO2]$ ,  $[\sum H_3PO4]$ ,  $[\sum Si(OH)_4]$ ,  $[\sum B(OH)_3]$ ,  $[\sum H_2SO_4]$ ,  $[\sum HF]$ , as well as water itself is calculated
- if the flag revelle=TRUE is set, the revelle factor (Zeebe and Wolf-Gladrow 2001) is calculated. item if the flag revelle=TRUE is set, all necessary quantities for the explicit "direct substitution approach" (DSA) to pH modelling as given in Hofmann, Meysman, Soetaert, and Middelburg (2008b) are calculated. These are the buffer factor (the partial derivative of [TA] with respect to  $[H^+]$ ) and the partial derivatives of [TA] with respect to the other total quantities. Furthermore, the partial derivatives of [TA] with respect to changes in the equilibrium constants  $(K^*)$ , multiplied with the partial derivatives of the equilibrium constants with respect to their variables needed for the DSA with time variable equilibrium constants as described in Hofmann, Meysman, Soetaert, and Middelburg (2008a) are calculated. Finally, the ionization fractions as defined by Stumm and Morgan (1996) and used in Hofmann, Middelburg, Soetaert, Wolf-Gladrow, and Meysman (2008c) are calculated for the full speciation.
- Input for aquaenv has to be supplied in standard SI units, the free proton pH scale and in molinity<sup>1</sup> (mol/kg-solution). Conversion of input parameters to this necessary units and pH scale can be done with the generic function convert.
- The information created with aquaenv is also supplied in standard SI units and in molinity. All elements of an object of class aquaenv of a certain unit or pH scale can be converted into other units or pH scales with the function convert as well.
- One can use input vectors of temperature T, salinity S or depth d for aquaenv to obtain vectors of all calculated information as function of the input vector. This can be visualized in a large variety of ways using the plot function specially defined for objects of type aquaenv.
- Objects of class aquaenv can be used in dynamic models to define the state of the
  system in each timestep of the numerical integration (done e.g. with deSolve). with the
  function aquaenv and the flag from.data.frame=TRUE it is possible to convert output
  of those dynamic models into objects of type aquaenv which allows the user to use the
  whole suite of visualisation tools that is provided by the function plot in AquaEnv.
- As mentioned above Hofmann et al. (2008b), Hofmann et al. (2008a), and Hofmann et al. (2008c) describe methods for an "explicit" pH modelling which allows for the quantification of the influences of kinetically modelled processes on the pH. Objects of type aquaenv provide all needed quantities (partial derivatives of [TA], ionization fractions, etc.) to employ both of those methods in dynamic models. Furthermore, AquaEnv provides the functionality to cumulatively plot the obtained influences on the pH.
- As an example of how to use the toolbox that is **AquaEnv**, two applications are provided

<sup>&</sup>lt;sup>1</sup>Note that it is not sufficient to give a gravimetric concentration in mol/kg since there is a substancial difference between mol/kg-H<sub>2</sub>O (molality) and mol/kg-solution (molinity).

- The function titration: creates theoretical titrations which can be used e.g. to create bjerrum plots, something that can also be done with the function plot in AquaEnv.
- − The function TAfit: a routine based on a method in DOE (1994) that makes use of that theoretical titration function and allows for determining total alkalinity ([TA]), the total dissolved inorganic carbon concentration ([ $\sum$ CO2]), as well as additionally the electrode standard potential (E<sub>0</sub>) and the first dissociation constant of the carbonate system ( $K_{CO_2}^*$ ) using the Levenberg-Marquart algorithm (least squares optimization procedure) as provided in **minpack.lm**.

# 2 The elements of an object of class aquaenv

The function aquaenv, the central function of **AquaEnv**, returns an object of class aquaenv. This object is a list of different elements which can be accesses with the \$\\$ character or with the [[]] operator

```
> test <- aquaenv(10, 35)
> test$Tc
> test[["Tc"]]
```

Maximally, i.e., if the enough input data is supplied to define the pH of the system and the flags speciation, dsa, and revelle are TRUE while the flag skeleton is FALSE, an object of class aquaenv contains the following elements

element	unit	explanation
$\mathrm{Tc}$	°C	temperature
$\operatorname{Tk}$	K	absolute temperature
S	"psu" (no unit)	salinity
Cl	%0	chlorinity
I	mol/kg-H <sub>2</sub> O	ionic strength
d	m	depth
hydroP	bar	hydrostatic pressure
density	$kg/m^3$	(seawater) density
SumCO2	mol/kg-soln	$[\sum CO_2]$ , total dissolved inorganic carbon concentration
SumNH4	mol/kg-soln	$[\sum NH_4^+]$ , total ammonium concentration
SumH2S	mol/kg-soln	$[\sum H_2S]$ , total sulfide concentration
SumHNO3	mol/kg-soln	$[\sum HNO_3]$ , total nitrate concentration
SumHNO2	mol/kg-soln	$[\sum HNO_2]$ , total nitrite concentration
SumH3PO4	mol/kg-soln	$[\sum H_3PO_4]$ , total phosphate concentration
SumSiOH4	mol/kg-soln	$[\sum Si(OH)_4]$ , total silicate concentration
SumBOH3	mol/kg-soln	$[\sum B(OH)_3]$ , total borates concentration
SumH2SO4	mol/kg-soln	$[\sum H_2SO_4]$ , total sulfate concentration
SumHF	mol/kg-soln	$[\sum HF]$ , total fluoride concentration
SumBr	mol/kg-soln	[\sum_HBr], total bromide concentration
ClConc	mol/kg-soln	[Cl <sup>-</sup> ], chloride concentration

Na	mol/kg-soln	[Na <sup>+</sup> ], sodium concentration
Mg	mol/kg-soln	$[Mg^{2+}]$ , magnesium concentration
Ca	mol/kg-soln	[Ca <sup>2+</sup> ], calcium concentration
K	mol/kg-soln	[K <sup>+</sup> ], potassium concentration
Sr	mol/kg-soln	[Sr <sup>2+</sup> ], strontium concentration
molal2molin	(mol/kg-soln)/(mol/kg-H2O)	concentration conversion factor: from molality to molinity
free2tot	-	pH conversion factor: free scale to total scale
free2sws	-	pH conversion factor: free scale to sawater scale
tot2free	-	pH conversion factor: total scale to free scale
tot2sws	-	pH conversion factor: total scale to seawater scale
sws2free	-	pH conversion factor: seawater scale to free scale
sws2tot	-	pH conversion factor: seawater scale to total scale
K0_CO2	mol/(kg-soln*atm)	Henry's constant for CO <sub>2</sub>
K0_O2	mol/(kg-soln*atm)	Henry's constant for O <sub>2</sub>
CO2_sat	mol/kg-soln	${\rm CO_2}$ saturation concentration at an atmospheric partial pressure/fugacity of Fugacity\$CO2
O2_sat	mol/kg-soln	${\rm O}_2$ saturation concentration at an atmospheric partial pressure/fugacity of Fugacity\$O2
K_W	$(\text{mol/kg-soln})^2$ , free pH scale	stoichiometric equilibrium ion product of
		$H_2O: K_W^* = [H^+][OH-]$
K_HSO4	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
		$K_{HSO_4^-}^* = [H^+][SO_4^{2-}]/[HSO_4^-]$
K_HF	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
	, , , ,	$K_{HF}^* = [H^+][F^-]/[HF]$
K_CO2	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
		$K_{CO_2}^* = [H^+][HCO_3^-]/[CO_2]$
K_HCO3	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
		$K_{HCO_3^-}^* = [H^+][CO_3^{2-}]/[HCO_3^-]$
K_BOH3	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
	, G , 1	$K_{B(OH)_3}^* = [H^+][B(OH)_4^-]/[B(OH)_3]$
K_NH4	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
	, , , ,	$K_{NH_4^+}^* = [H^+][NH_3]/[NH_4^+]$
K_H2S	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
		$K_{H_2S}^* = [H^+][HS^-]/[H_2S]$
K_H3PO4	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
		$K_{H_3PO_4}^* = [H^+][H_2PO_4^-]/[H_3PO_4]$
K_H2PO4	$\mathrm{mol/kg} ext{-}\mathrm{soln},\;\mathrm{free}\;\mathrm{pH}\;\mathrm{scale}$	stoichiometric equilibrium constant
		$K_{H_2PO_4^-}^* = [H^+][HPO_4^{2-}]/[H_2PO_4^-]$
K_HPO4	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
		$K_{HPO_4^{2-}}^* = [H^+][PO_4^{3-}]/[HPO_4^{2-}]$
K_SiOH4	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
	, G , F Source	$K_{Si(OH)_4}^* = [H^+][SiO(OH)_3^-]/[Si(OH)_4]$
K_SiOOH3	mol/kg-soln, free pH scale	stoichiometric equilibrium constant
3-2 22	, pri som	$K_{SiO(OH)_3}^* = [H^+][SiO_2(OH)_2^{2-}]/[SiO(OH)_3^{-}]$
		$SiO(OH)_3$ $I = 2 \cdot 2 \cdot 2 \cdot 1 / 1 \cdot 2 \cdot 3 \cdot 1$

K_HNO3 $K_{HNO_2} = [H^+][NO_2^-]/[HNO_2]$ approximate value for equilibrium constant $K_{HNO_3}^* = [H^+][NO_3^-]/[HNO_3]$ approximate value for equilibrium constant $K_{HNO_3}^* = [H^+][NO_3^-]/[HNO_3]$ approximate value for equilibrium constant $K_{H2SO_4}^* = [H^+][HSO_4^-]/[H_2SO_4]$ approximate value for equilibrium constant $K_{HS^-}^* = [H^+][S^2^-]/[HS^-]$ stoichiometric equilibrium solubility product of
K_H2SO4 mol/kg-soln; mol/kg-H2O; mol/l approximate value for equilibrium constant $K_{\rm H2SO_4}^* = [{\rm H^+}][{\rm HSO_4^-}]/[{\rm H_2SO_4}]$ approximate value for equilibrium constant $K_{\rm HS^-}^* = [{\rm H^+}][{\rm S^2^-}]/[{\rm HS^-}]$
K_HS $ \begin{array}{c} K_{\rm H_2SO_4} = [{\rm H^+}][{\rm HSO_4^-}]/[{\rm H_2SO_4}] \\ {\rm approximate\ value\ for\ equilibrium\ constant} \\ K_{\rm HS^-}^* = [{\rm H^+}][{\rm S^2^-}]/[{\rm HS^-}] \end{array} $
K_HS
$K_{HS^-}^* = [H^+][S^{2-}]/[HS^-]$
calcite
$Ksp_{cal}^* = [Ca^{2+}][CO_3^{2-}]$
Ksp_aragonite $(\text{mol/kg-soln})^2$ stoichiometric equilibrium solubility product of aragonite
$Ksp_{ara}^* = [Ca^{2+}][CO_3^{2-}]$
TA mol/kg-soln [TA], total alkalinity
pH -, free scale pH
pCO2 atm, partial pressure (fugacity) of CO <sub>2</sub> in the water
$CO2$ $mol/kg$ -soln $[CO_2]$
$\mathrm{HCO3}$ $\mathrm{mol/kg\text{-}soln}$ $\mathrm{[HCO_3^-]}$
$CO3$ $mol/kg$ -soln $[CO_3^{2-}]$
BOH3 $\text{mol/kg-soln}$ $[B(OH)_3]$
BOH4 $\text{mol/kg-soln}$ $[B(OH)_4^-]$
$OH   mol/kg$ -soln $[OH^-]$
$H3PO4$ $mol/kg$ -soln $[H_3PO_4]$
$H2PO4$ $mol/kg$ -soln $[H2PO_4^-]$
$\mathrm{HPO4}$ $\mathrm{mol/kg\text{-}soln}$ $\mathrm{[HPO_4^{2-}]}$
PO4 $\text{mol/kg-soln}$ $[PO_4^{3-}]$
SiOH4 $\text{mol/kg-soln}$ $\left[\text{Si(OH)}_4\right]$
SiOOH3 $\text{mol/kg-soln}$ $\left[\text{SiO(OH)}_3^-\right]$
SiO2OH2 $\text{mol/kg-soln}$ $\left[\text{SiO}_2(\text{OH})_2^{2-}\right]$
$H2S$ $mol/kg$ -soln $[H_2S]$
$HS$ $mol/kg$ -soln $[HS^-]$
S2min $mol/kg$ -soln $[S^{2-}]$
$NH4$ $mol/kg$ -soln $[NH_4^+]$
NH3 $mol/kg$ -soln $[NH_3]$
$H2SO4$ $mol/kg$ -soln $[H_2SO_4]$
$HSO4$ $mol/kg$ -soln $[HSO_4^-]$
SO4 $\operatorname{mol/kg-soln}$ $[SO_4^{2-}]$
HF mol/kg-soln [HF]
$[F^-]$
$HNO3$ $mol/kg$ -soln $[HNO_3]$
NO3 $\text{mol/kg-soln}$ $[\text{NO}_3^-]$
$HNO2$ $mol/kg$ -soln $[HNO_2]$
$NO2$ $mol/kg$ -soln $[NO_2^-]$
omega_calcite - saturation state $\Omega$ with respect to calcite
omega_aragonite - saturation state $\Omega$ with respect to a ragonite

revelle	-	Revelle factor
c1	-	ionization fraction $c_1 = [CO_2]/[\sum CO_2]$
c2	-	ionization fraction $c_2 = [HCO_3^-]/[\sum CO_2]$
c3	-	ionization fraction $c_3 = [CO_3^{2-}]/[\sum CO_2]$
${\rm dTAdSumCO2}$	-	$\frac{\partial[\text{TA}]}{[\partial \sum \text{CO}_2]}$
		with $[TA] = f([H^+], [\sum CO_2],)$
b1	_	ionization fraction $b_1 = [B(OH)_3]/[\sum B(OH)_3]$
b2	_	ionization fraction $b_2 = [B(OH)_4^-]/[\sum B(OH)_3]$
dTAdSumBOH3	-	$\frac{\partial [TA]}{[\partial \sum B(OH)_3]}$
		with $[TA] = f([H^+], [\sum CO_2],)$
so1	-	ionization fraction $so_1 = [H_2SO_4]/[\sum H_2SO_4]$
so2	-	ionization fraction $so_2 = [HSO_4^-]/[\sum H_2SO_4]$
so3	-	ionization fraction $so_3 = [SO_4^{2-}]/[\sum H_2SO_4]$
dTAdSumH2SO4	-	$ \frac{\partial [\text{TA}]}{[\partial \sum \text{H}_2 \text{SO}_4]} $
		with $[TA] = f([H^+], [\sum CO_2],)$
f1	-	ionization fraction $f_1 = [HF]/[\sum HF]$
f2	-	ionization fraction $f_1 = [F^-]/[\sum HF]$
dTAdSumHF	-	$ \frac{\partial [\text{TA}]}{[\partial \sum \text{HF}]} $
		with $[TA] = f([H^+], [\sum CO_2],)$
dTAdH	-	$\frac{\partial[TA]}{[\partial[H^+]]}$ : buffer factor
		with $[TA] = f([H^+], [\sum CO_2],)$
$\mathrm{dTAdKdKdS}$	-	$\sum_{i} \frac{\partial [TA]}{\partial K_{i}^{*}} \frac{\partial K_{i}^{*}}{\partial S}$
		with $[TA] = f([H^+], [\sum CO_2],, K_i^*)$
dTAdKdKdT	-	$\sum_{i} \frac{\partial [TA]}{\partial K_{i}^{*}} \frac{\partial K_{i}^{*}}{\partial T}$
		with $[TA] = f([H^+], [\sum CO_2],, K_i^*)$
dTAdKdKdd	-	$\sum_{i} \frac{\partial [TA]}{\partial K_{i}^{*}} \frac{\partial K_{i}^{*}}{\partial d}$
		with $[TA] = f([H^+], [\sum CO_2],, K_i^*)$
dTAdKdKdSumH2SO4	-	$\sum_{i} \frac{\partial [TA]}{\partial K_{i}^{*}} \frac{\partial K_{i}^{*}}{\partial [\sum H_{2}SO_{4}]}$
		with $[TA] = f([H^+], [\sum CO_2],, K_i^*)$
${\rm dTAdKdKdSumHF}$	-	$\sum_{i} \frac{\partial [TA]}{\partial K_{i}^{*}} \frac{\partial K_{i}^{*}}{\partial [\sum HF]}$
		with $[TA] = f([H^+], [\sum CO_2],, K_i^*)$
	ı	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

For elements that are calculated according to certain literature references, those references are given in appendix A.

# 3 Using AquaEnv

# 3.1 Basic features

# 3.1.1 calling the "K" functions directly

The elements K\_CO2, K\_HCO3, K\_BOH3, K\_W, K\_HSO4, K\_HF, K\_NH4, K\_H2S, K\_H3PO4, K\_H2PO4, K\_HPO4, K\_SiOH4, K\_SiOH3, KO\_CO2, KO\_O2, Ksp\_aragonite, and Ksp\_calcite can be calculated directly, witout creating an object of class *aquaenv*. This is done via functions that bear the same as those elements

```
> K_C02(15, 30)
> K0_C02(15, 30)
> Ksp_calcite(15, 30, 100)
```

## 3.1.2 Minimal aquaenv definition

Minimally, an object of class *aquaenv* can be defined with just a temperature and salinity value

```
> ae <- aquaenv(Tc = 15, S = 30)
> ae$K_CO2
```

Optionally, a mean depth can be given. As in the above case, the returned object of class aquaenv then contains a standard set of elements as shown by the names command.

```
> ae <- aquaenv(Tc = 15, S = 30, d = 10)
> ae$Ksp_calcite
> names(ae)
```

A minimal set of elements in an object of class aquaenv can be obtained by setting the flag skeleton to TRUE

```
> ae <- aquaenv(Tc = 15, S = 30, d = 10, skeleton = TRUE) > names(ae)
```

#### 3.1.3 Defining the complete aquaenv system in different ways

If enough information is given to define a complete speciation, i.e. either one of the pairs SumCO2 and pH, SumCO2 and TA, SumCO2 and CO2, or SumCO2 and pCO2, a full aquaenv system can be defined.

```
> Tc <- 15
> S <- 30
> d <- 10
> SumCO2 <- 0.002
> pH <- 8
> TA <- 0.002140323
> pCO2 <- 0.000533576
> CO2 <- 2.055419e-05
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, pH = pH)
> ae$TA
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, TA = TA)
> ae$pH
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, CO2 = CO2)
> ae$pH
> names(ae)
```

As seen above, a full speciation is calculated along with the pH or total alkalinity respectively. If only pH or total alkalinity is needed, the calculation of the full speciation can be toggled off. Furthermore the flag skeleton also works for a full system.

```
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, pH = pH, speciation = FALSE)
> names(ae)
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, pH = pH, speciation = FALSE,
+ skeleton = TRUE)
> names(ae)
```

Furtermore all the quantities needed for the explicit pH modelling approaches as given in Hofmann *et al.* (2008b) and Hofmann *et al.* (2008c) can be calculated by setting the flag dsa to TRUE. The revelle factor can be calculated using the flag revelle.

```
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, pCO2 = pCO2,
+ dsa = TRUE, revelle = TRUE)
> ae$dTAdH
> ae$revelle
```

If an ambivalent situation is created because the user enters too much information, an error message is displayed

```
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, CO2 = CO2, pCO2 = pCO2)
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, pH = pH, TA = TA)
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, pH = pH, CO2 = CO2)
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, pH = pH, pCO2 = pCO2)
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, TA = TA, CO2 = CO2)
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, TA = TA, pCO2 = pCO2)
> ae <- aquaenv(Tc, S, d, SumCO2 = SumCO2, TA = TA, pCO2 = pCO2)
```

# 3.1.4 Calculating $[\sum CO_2]$

 $[\sum CO_2]$  can be calculated by giving a constant pair of either pH and CO2, pH and pCO2, pH and TA, TA and CO2, or TA and pCO2

```
> pC02 <- 0.0006952296
> C02 <- 2.678134e-05
> pH <- 7.888569
> TA <- 0.0021
> Tc <- 15
> S <- 30
> d <- 10
> ae <- aquaenv(Tc, S, d, SumCO2 = NULL, pH = pH, CO2 = CO2)
> ae$SumCO2
> ae <- aquaenv(Tc, S, d, SumCO2 = NULL, pH = pH, pCO2 = pCO2)
> ae$SumCO2
> ae <- aquaenv(Tc, S, d, SumCO2 = NULL, pH = pH, TA = TA)
> ae$SumCO2
> ae <- aquaenv(Tc, S, d, SumCO2 = NULL, pH = pH, TA = TA)
> ae$SumCO2
> ae <- aquaenv(Tc, S, d, SumCO2 = NULL, pH = pH, TA = TA)</pre>
```

```
> ae$SumCO2
> ae <- aquaenv(Tc, S, d, SumCO2 = NULL, TA = TA, pCO2 = pCO2)
> ae$SumCO2
```

## 3.1.5 Cloning an object of class aquaenv

It is possible to clone an obect of class aquaenv, either 1 to 1 or with different pH, TA, or K\_CO2

```
> Tc <- 15
> S <- 30
> SumCO2 <- 0.002
> TA <- 0.00214
> ae <- aquaenv(Tc, S, SumCO2 = SumCO2, TA = TA)
> ae$pH
> ae1 <- aquaenv(ae = ae)
> ae1$pH
> pH <- 9
> ae2 <- aquaenv(ae = ae, pH = pH)
> ae2$TA
> TA <- 0.002
> ae3 <- aquaenv(ae = ae, TA = TA)
> ae3$pH
> K_CO2 <- 1e-06
> ae4 <- aquaenv(ae = ae, k_co2 = K_CO2)
> ae4$pH
```

Note that K\_CO2 as an input variable is in lower cases!

# 3.1.6 Preparing input variables

Input variables for the function aquaenv need to be in mol/kg-solution and on the free pH scale. Data in other concentration units or pH scales can be converted using the function convert.

# 3.1.7 Vectors as input variables

One of the input variables for the function aquaenv may be a vector. All the other input variables are then assumed to be constant. The elements of the resulting two dimensional object of class aquaenv are then vectors containing the elements as a function of the input variable for which a vector is given.

```
> SumC02 <- 0.002
> pH <- 8
> Tc <- 1:15
> S <- 30
> d <- 10
> ae <- aquaenv(Tc, S, d, SumC02 = SumC02, pH = pH, revelle = TRUE)
> ae$revelle
```

The plot function plots all elements of the respective object of class aquaenv. This, however, Anight distributed the class aquaenvial of the respective object of class aquaenv. This, however, Anight distributed the class aquaenvial of the respective distributed the control of the control o

```
> plot(ae, xval = Tc, xlab = "T/(deg\ C)", what = c("pH", + "CO2", "HCO3", "CO3"), newdevice = FALSE, mfrow = c(1, 4))
```

The following chunks of example code show other possible definitions of objects of class aquaenv with vectors as input variables.

Interesting to note is that also, e.g., SumCO2, TA, pH and SumNH4 can be vectors

```
> ae <- aguaenv(10, 20, SumCO2 = seq(0.001, 0.002, 1e-04),
      TA = 0.002)
> plot(ae, xval = ae$SumCO2, xlab = "SumCO2/(mol/kg-soln)",
      what = c("pH", "CO2", "HCO3", "CO3"), newdevice = FALSE,
      mfrow = c(1, 4)
> ae <- aquaenv(10, 20, SumCO2 = 0.002, TA = seq(0.001,
      0.002, 1e-04))
> plot(ae, xval = ae$TA, xlab = "TA/(mol/kg-soln)", what = c("pH",
      "CO2", "HCO3", "CO3"), newdevice = FALSE, mfrow = c(1, 
      4))
> ae <- aquaenv(10, 20, SumCO2 = 0.002, pH = seq(8, 8.2,
      0.001))
> plot(ae, xval = aepH, xlab = "pH (free scale)", what = c("pH",
      "CO2", "HCO3", "CO3"), newdevice = FALSE, mfrow = c(1,
      4))
> ae <- aquaenv(10, 20, SumCO2 = 0.002, SumNH4 = seq(1e-04,
      2e-04, 1e-05), pH = 8)
> ae$NH3
```

# 3.1.8 Calculating $[\sum CO_2]$ from input vectors

The functionality of calculating SumCO2 can also be used together with vectors as input variables.

```
> ae <- aquaenv(Tc = 11:15, S = 30, SumCO2 = NULL, pH = pH, + CO2 = CO2, revelle = TRUE, dsa = TRUE) > ae$SumCO2
```

Two further examples

#### 3.1.9 Conversion from and to a dataframe

Objects of class aquaenv can be converted to an R data.frame to further post-process them with standard R means. Similarly, R data.frames can be converted to objects of class aquaenv to use the plotting facilities that exist for objects of class aquaenv. This can be helpful for plotting output of a dynamic model run, e.g. from R package **deSolve**, and will be shown later in this document.

```
> aedataframe <- as.data.frame(ae)
> aetest <- aquaenv(ae = aedataframe, from.data.frame = TRUE)</pre>
```

#### 3.1.10 Converting elements in an obect of class aquaenv

Elements of an object of class *aquaenv* are calculated in, e.g., the concentration unit mol/kg-solution (molinity). The function **convert** can be used to convert all elements in an object of class *aquaenv* that share a common attribute, e.g. the unit.

# 3.1.11 Quantities needed for explicit pH modelling

As already mentioned above, the quantities needed for the explicit pH modelling approach (direct substitution approach - DSA) as presented by Hofmann *et al.* (2008b) can be calculated with the function aquaenv by setting the flag dsa.

```
> ae <- aquaenv(Tc = 15, S = 30, d = 10, SumCO2 = 0.002, + pH = 8, dsa = TRUE, revelle = TRUE)
```

This command calculated the buffer factor and the partial derivatives of [TA] with respect to other summed quantities referred to in Hofmann *et al.* (2008b)

```
> ae$dTAdH
> ae$dTAdSumCO2
```

as well the sums partial derivatives of [TA] with respect to the equilibrium constants ( $K^*$ 's) multiplied with the partial derivatives of the respective equilibrium constant with one of their variables (i.e., S, T, d, SumH2SO4, od SumHF) as introduced in Hofmann *et al.* (2008a).

```
ae$dTAdKdKdSae$dTAdKdKdSumH2SO4
```

Furthermore the ionization fractions used for the pH dependent fractional stoichiometric pH modelling approach described in Hofmann et al. (2008c) are calculated as well

```
> ae$c1
```

# 3.2 The plot.aquaenv function

In the previous sections, the plot function has been introduced. What actually is called if the first element of the arguments list of plot is an object of type aquaenv is the function plot.aquaenv. This is a multifunctional tool to visualize information contained in an object of class aquaenv. For the convenience of the users, plot.aquaenv combines the call of standard R plotting functions and the previous call of the function par to set parameters like mfrow, mar, etc. as well as the opening of a plotting device with a certain size. As already shown above, setting the flag newdevice to FALSE suppresses the opening of a new plotting device (this feature is needed here to create a plot that will be woven into the LATEX document by Sweave).

For example

Furthermore the parameter device can be specified which allows the user to write the plots to .eps and .pdf files. The parameter filename can be used to specify a filename other than the default filename "aquaenv".

These features make the function plot.aquaenv different from standard R plotting functions. However, if the flags newdevice and setpar are set to FALSE, plot.aquaenv behaves like a "normal" R plotting function

```
> par(mfrow = c(1, 2))
> plot(ae, xval = 20:30, xlab = "S", what = "K_CO2", lwd = 3,
+ col = "red", newdevice = FALSE, setpar = FALSE)
> plot(ae, xval = 20:30, xlab = "S", what = "K_HCO3", cex = 3,
+ type = "b", col = "blue", newdevice = FALSE, setpar = FALSE)
```

Furthermore, the function plot.aquaenv can be used to create "cumulative" plots and "Bjerrum" plots. This will be explained in some of the following sections.

## 3.3 Using objects of class aquaenv in dynamic models

#### 3.3.1 Ordinary dynamic models

It is convenient to use objects of class *aquaenv* in a dynamic model, e.g. solved using the R package **deSolve**. This can be illustrated with an example. (For information about how to set up a dynamic model with **deSolve**, consult the documentation of **deSolve**).

```
> require(deSolve)
```

A list of parameters is specified

```
> parameters <- list(S = 25, Tc_min = 5, Tc_max = 25, d = 10,
+ k = 0.4, r0x = 3e-07, rNitri = 2e-07, rPP = 6e-06,
+ ksDINPP = 1e-06, ksNH4PP = 1e-06, D = 0.1, 02_io = 0.000296,
+ NO3_io = 3.5e-05, SumNH4_io = 8e-06, SumCO2_io = 0.00232,
+ TA_io = 0.002435, C_Nratio = 8, a = 30, b = 50, modeltime = 100)</pre>
```

A model function is defined which will be executed every timestep of the numerical integration. An object of class *aquaenv* is created in each timestep, some of its elements are used to calculated kinetic rate expressions and the whole object is returned as output.

```
> Waddenzeebox <- function(timestep, currentstate, parameters) {
      with(as.list(c(currentstate, parameters)), {
          Tc \leftarrow c(seq(Tc_min, Tc_max, (Tc_max - Tc_min)/(modeltime/2)),
               seq(Tc_max, Tc_min, -(Tc_max - Tc_min)/(modeltime/2)))[[round(timestep) +
               1]]
          ae \leftarrow aquaenv(Tc = Tc, S = S, SumCO2 = SumCO2,
               SumNH4 = SumNH4, TA = TA)
          ECO2 \leftarrow k * (ae$CO2_sat - ae$CO2)
          E02 <- k * (ae$02_sat - 02)
          TO2 <- D * (O2_io - O2)
          TNO3 <- D * (NO3_io - NO3)
          TSumNH4 <- D * (SumNH4_io - SumNH4)
          TTA \leftarrow D * (TA_{io} - TA)
          TSumCO2 \leftarrow D * (SumCO2_io - SumCO2)
          RNit <- rNitri
          ROx <- rOx
          ROxCarbon <- ROx * C_Nratio
          pNH4PP <- 0
          RPP <- 0
          if ((timestep > a) && (timestep < b)) {
              RPP \leftarrow rPP * ((SumNH4 + NO3)/(ksDINPP + (SumNH4 +
                   NO3)))
               pNH4PP <- 1 - (ksNH4PP/(ksNH4PP + SumNH4))
          else {
              RPP <- 0
```

```
RPPCarbon <- RPP * C_Nratio</pre>
+
          d02 \leftarrow T02 + E02 - R0xCarbon - 2 * RNit + (2 -
               2 * pNH4PP) * RPP + RPPCarbon
          dNO3 \leftarrow TNO3 + RNit - (1 - pNH4PP) * RPP
          dSumCO2 <- TSumCO2 + ECO2 + ROxCarbon - RPPCarbon
          dSumNH4 <- TSumNH4 + ROx - RNit - pNH4PP * RPP
          dTA \leftarrow TTA + ROx - 2 * RNit - (2 * pNH4PP - 1) *
               RPP
          ratesofchanges <- c(d02, dN03, dSumNH4, dSumC02,
               dTA)
           transport \leftarrow c(T02 = T02, TN03 = TN03, TSumNH4 = TSumNH4,
               TTA = TTA, TSumCO2 = TSumCO2)
          airseaexchange \leftarrow c(ECO2 = ECO2, EO2 = EO2)
          return(list(ratesofchanges, list(transport, airseaexchange,
               ae)))
      })
+ }
```

The model is solved

and the output can be plotted in the same way as a two dimensional object of class aquaenv by converting it to an object of class aquaenv using the from.data.frame flag of the function aquaenv

```
> plot(aquaenv(ae = output, from.data.frame = TRUE), xval = outputtime,
+ xlab = "time/d", mfrow = c(10, 10), newdevice = FALSE)
```

## 3.3.2 Models using the explicit pH modelling approach

**3.3.2.1** In one single model Since an object of class *aquaenv* can contain all quantities necessary to employ the explicit pH modelling approaches as introduced by Hofmann *et al.* (2008b,a,c), they can be readily used in an explicit pH model.

As an example, we give a model that calculates the pH in the "classical" way in every timestep using aquaenv, also employs the explicit pH modelling approach (direct substitution approach - DSA) given in Hofmann et al. (2008b) and additionally employs fractional stoichiometry as given in Hofmann et al. (2008c). The pH evolution is thus calculated in three different ways

which allows comparing the three values for consistency.

Again, a list of parameters is defined

```
> parameters <- list(S = 25, Tc = 15, d = 10, k = 0.4,
+ r0x = 3e-07, rNitri = 2e-07, rPP = 6e-07, ksSumNH4 = 1e-06,
+ D = 0.1, 02_io = 0.000296, NO3_io = 3.5e-05, SumNH4_io = 8e-06,
+ SumCO2_io = 0.00232, TA_io = 0.002435, C_Nratio = 8,
+ a = 30, b = 50, modeltime = 100)
```

And a model function is defined. Again, an object of class *aquaenv* is created in each timestep and respective elements are used.

```
> boxmodel <- function(timestep, currentstate, parameters) {
      with(as.list(c(currentstate, parameters)), {
           ae \leftarrow aquaenv(Tc = Tc, S = S, SumCO2 = SumCO2,
               SumNH4 = SumNH4, TA = TA, dsa = TRUE)
          ECO2 \leftarrow k * (ae$CO2\_sat - ae$CO2)
          E02 \leftarrow k * (ae$02_sat - 02)
          RNit <- rNitri
          ROx <- rOx
          if ((timestep > a) && (timestep < b)) {
               RPP <- rPP * (SumNH4/(ksSumNH4 + SumNH4))</pre>
           else {
               RPP <- 0
           d02 <- E02 - C_Nratio * ROx - 2 * RNit + C_Nratio *</pre>
               RPP
           dNO3 <- RNit
           dSumCO2 <- ECO2 + C_Nratio * ROx - C_Nratio *
               RPP
           dSumNH4 \leftarrow ROx - RNit - RPP
           dTA \leftarrow ROx - 2 * RNit - RPP
           dH <- (dTA - (dSumCO2 * ae$dTAdSumCO2 + dSumNH4 *
               ae$dTAdSumNH4))/ae$dTAdH
          DSApH \leftarrow -log10(H)
          rhoHECO2 <- ae$c2 + 2 * ae$c3
          rhoHRNit <-1 + ae$n1
           rhoHROx \leftarrow C_Nratio * (ae$c2 + 2 * ae$c3) - ae$n1
           rhoHRPP \leftarrow -(C_Nratio * (ae$c2 + 2 * ae$c3)) +
               ae$n1
          dH_ECO2 <- rhoHECO2 * ECO2/(-ae$dTAdH)</pre>
           dH_RNit <- rhoHRNit * RNit/(-ae$dTAdH)</pre>
           dH_ROx \leftarrow rhoHROx * ROx/(-ae$dTAdH)
           dH_RPP <- rhoHRPP * RPP/(-ae$dTAdH)</pre>
           dH_stoich <- dH_ECO2 + dH_RNit + dH_ROx + dH_RPP</pre>
```

```
# DSAstoichpH <- -log10(H_stoich)
# ratesofchanges <- c(dO2, dNO3, dSumNH4, dSumCO2,
# dTA, dH, dH_stoich)
# processrates <- c(ECO2 = ECO2, EO2 = EO2, RNit = RNit,
# ROx = ROx, RPP = RPP)
# DSA <- c(DSApH = DSApH, rhoHECO2 = rhoHECO2,
# rhoHRNit = rhoHRNit, rhoHROx = rhoHROx, rhoHRPP = rhoHRPP,
# dH_ECO2 = dH_ECO2, dH_RNit = dH_RNit, dH_ROx = dH_ROx,
# dH_RPP = dH_RPP, DSAstoichpH = DSAstoichpH)
# return(list(ratesofchanges, list(processrates,
# DSA, ae)))
# })
# })</pre>
```

The model is solved

and output can be plotted. Again using plot.aquaenv. Note that here the parameter what is used.

```
> what <- c("SumCO2", "TA", "SumNH4", "NO3", "ECO2", "EO2",
+ "RNit", "ROx", "RPP", "dTAdH", "dTAdSumCO2", "dTAdSumNH4",
+ "c1", "c2", "c3", "n1", "n2", "rhoHECO2", "rhoHRNit",
+ "rhoHROx", "rhoHRPP", "dH_ECO2", "dH_RNit", "dH_ROx",
+ "dH_RPP", "pH", "DSApH", "DSAstoichpH")
> plot(aquaenv(ae = output, from.data.frame = TRUE), xval = output$time,
+ what = what, xlab = "time/d", mfrow = c(6, 5), size = c(20,
+ 13), newdevice = FALSE)
```

Here, the cumulative plotting functionality of plot.aquaenv can be employed as well to visualize the influences of the different kinetically modelled processes on [H<sup>+</sup>].

```
> what <- c("dH_ECO2", "dH_RNit", "dH_ROx", "dH_RPP")
> plot(aquaenv(ae = output, from.data.frame = TRUE), xval = output$time,
+ what = what, xlab = "time/d", size = c(7, 5), ylab = "mol-H/(kg-soln*d)",
+ legendposition = "bottomright", cumulative = TRUE,
+ newdevice = FALSE)
```

Finally, the pH values calculated with the three different methods can be plotted in one single graph to see that they are identical, i.e. the three methods of pH calculation are consistent with each other

```
> ylim <- range(output$DSApH, output$DSAstoichpH, output$pH)
> plot(output$DSApH, ylim = ylim, type = "l", xlab = "time/d",
+     ylab = "pH (free scale)")
> par(new = TRUE)
> plot(output$DSApH, ylim = ylim, type = "l", col = "red",
+     xlab = "", ylab = "")
> par(new = TRUE)
> plot(output$DSAstoichpH, ylim = ylim, type = "l", col = "blue",
+     xlab = "", ylab = "")
```

#### 3.3.2.2 In three separate models

## **3.3.2.2.1** The implicit pH modelling approach A list of parameters

```
> parameters <- list(S = 35, Tc = 15, SumCO2_t0 = 0.002,
      TA_t0 = 0.0022, kc = 0.5, kp = 1e-06, n = 2, modeltime = 20,
      outputsteps = 100)
The model function
> boxmodel <- function(timestep, currentstate, parameters) {</pre>
      with(as.list(c(currentstate, parameters)), {
           ae \leftarrow aquaenv(Tc = Tc, S = S, SumCO2 = SumCO2,
               TA = TA, SumSiOH4 = 0, SumBOH3 = 0, SumH2SO4 = 0,
               SumHF = 0
          Rc \leftarrow kc * ((ae$CO2_sat) - (ae$CO2))
          Rp <- kp * (1 - ae$omega_calcite)^n</pre>
          dSumCO2 <- Rc - Rp
          dTA \leftarrow -2 * Rp
          ratesofchanges <- c(dSumCO2, dTA)
          processrates \leftarrow c(Rc = Rc, Rp = Rp)
          return(list(ratesofchanges, list(processrates,
               ae)))
      })
+ }
Solving the model
> with(as.list(parameters), {
      initialstate <<- c(SumCO2 = SumCO2_t0, TA = TA_t0)</pre>
      times <<- seq(0, modeltime, (modeltime/outputsteps))</pre>
      output <<- as.data.frame(vode(initialstate, times,</pre>
           boxmodel, parameters, hmax = 1))
+ })
```

Visualizing the output

```
> what <- c("SumCO2", "TA", "Rc", "Rp", "omega_calcite",
      "pH")
> plot(aquaenv(ae = output, from.data.frame = TRUE), xval = output$time,
      xlab = "", mfrow = c(2, 3), size = c(12, 5), what = what,
      newdevice = FALSE)
3.3.2.2.2
          The explicit pH modelling approach A list of parameters
> parameters <- list(S = 35, Tc = 15, SumCO2_t0 = 0.002,
      TA_t0 = 0.0022, kc = 0.5, kp = 1e-06, n = 2, modeltime = 20,
      outputsteps = 100)
The model function
> boxmodel <- function(timestep, currentstate, parameters) {</pre>
      with(as.list(c(currentstate, parameters)), {
          ae \leftarrow aquaenv(Tc = Tc, S = S, SumCO2 = SumCO2,
               pH = -log10(H), SumSiOH4 = 0, SumBOH3 = 0,
               SumH2SO4 = 0, SumHF = 0, dsa = TRUE)
          Rc \leftarrow kc * ((ae$CO2_sat) - (ae$CO2))
          Rp <- kp * (1 - ae$omega_calcite)^n</pre>
          dSumCO2 <- Rc - Rp
          dHRc <- (-(ae$dTAdSumCO2 * Rc))/ae$dTAdH
          dHRp \leftarrow (-2 * Rp - (ae dTAdSumCO2 * (-Rp)))/ae dTAdH
          dH <- dHRc + dHRp
          ratesofchanges <- c(dSumCO2, dH)</pre>
          processrates \leftarrow c(Rc = Rc, Rp = Rp)
          outputvars <- c(dHRc = dHRc, dHRp = dHRp)
          return(list(ratesofchanges, list(processrates,
               outputvars, ae)))
      })
+ }
Solving the model
> with(as.list(parameters), {
      aetmp \leftarrow aquaenv(Tc = Tc, S = S, SumCO2 = SumCO2\_tO,
           TA = TA_t0, SumSiOH4 = 0, SumBOH3 = 0, SumH2SO4 = 0,
           SumHF = 0
      H_t0 \leftarrow 10^(-aetmp$pH)
      initialstate <<- c(SumCO2 = SumCO2_t0, H = H_t0)</pre>
      times <<- seq(0, modeltime, (modeltime/outputsteps))</pre>
      output <<- as.data.frame(vode(initialstate, times,</pre>
           boxmodel, parameters, hmax = 1))
+ })
```

Visualizing the output

```
> what <- c("SumCO2", "TA", "Rc", "Rp", "omega_calcite",
      "pH", "dHRc", "dHRp")
> plot(aquaenv(ae = output, from.data.frame = TRUE), xval = output$time,
      xlab = "time/d", mfrow = c(3, 3), size = c(15, 10),
      what = what, newdevice = FALSE)
Cumulatively plotting the influences of the two processes on the pH
> what <- c("dHRc", "dHRp")</pre>
> plot(aquaenv(ae = output, from.data.frame = TRUE), xval = output$time,
      xlab = "time/d", what = what, ylab = "mol-H/(kg-soln*d)",
      legendposition = "topright", cumulative = TRUE, newdevice = FALSE)
3.3.2.2.3 The fractional stoichiometric approach A list of parameters
> parameters <- list(S = 35, Tc = 15, SumCO2_t0 = 0.002,
      TA_t0 = 0.0022, kc = 0.5, kp = 1e-06, n = 2, modeltime = 20,
      outputsteps = 100)
The model function
> boxmodel <- function(timestep, currentstate, parameters) {
      with(as.list(c(currentstate, parameters)), {
          ae \leftarrow aquaenv(Tc = Tc, S = S, SumCO2 = SumCO2,
              pH = -log10(H), SumSiOH4 = 0, SumBOH3 = 0,
              SumH2SO4 = 0, SumHF = 0, dsa = TRUE)
          Rc <- kc * ((ae$CO2_sat) - (ae$CO2))</pre>
          Rp <- kp * (1 - ae$omega_calcite)^n</pre>
          dSumCO2 <- Rc - Rp
          rhoc <- ae$c2 + 2 * ae$c3
          rhop <- 2 * ae$c1 + ae$c2
          dHRc <- rhoc * Rc/(-ae$dTAdH)
          dHRp <- rhop * Rp/(-ae$dTAdH)</pre>
          dH <- dHRc + dHRp
          ratesofchanges <- c(dSumCO2, dH)</pre>
          processrates \leftarrow c(Rc = Rc, Rp = Rp)
          outputvars <- c(dHRc = dHRc, dHRp = dHRp, rhoc = rhoc,
              rhop = rhop)
          return(list(ratesofchanges, list(processrates,
              outputvars, ae)))
      })
```

Solving the model

+ }

```
> with(as.list(parameters), {
      aetmp <- aquaenv(Tc = Tc, S = S, SumCO2 = SumCO2_t0,</pre>
          TA = TA_t0, SumSiOH4 = 0, SumBOH3 = 0, SumH2SO4 = 0,
          SumHF = 0
      H_t0 <- 10^{-aetmp$pH}
      initialstate <<- c(SumCO2 = SumCO2_t0, H = H_t0)
      times <<- seq(0, modeltime, (modeltime/outputsteps))</pre>
      output <<- as.data.frame(vode(initialstate, times,</pre>
          boxmodel, parameters, hmax = 1))
+ })
Visualizing the output
> what <- c("SumCO2", "TA", "Rc", "Rp", "omega_calcite",
      "pH", "dHRc", "dHRp", "rhoc", "rhop")
> plot(aquaenv(ae = output, from.data.frame = TRUE), xval = output$time,
      xlab = "time/d", mfrow = c(3, 4), size = c(15, 10),
      what = what, newdevice = FALSE)
```

#### 3.4 Titration simulation: the function titration

With the function **titration AquaEnv** provides a powerful tool to simulate titrations. A two dimensional object of class *aquaenv* will be created where the second dimension is the amount of titrant added. For this purpose, three extra elements are added to the *aquaenv* object

${f element}$	$\mathbf{unit}$	explanation		
delta_conc_titrant	mol/kg-solution	the offset in concentration of the titrant that is caused by adding the titrant to the sample		
$delta\_mass\_titrant$	kg	the amount of mass of titrant solution added		
$delta\_moles\_titrant$	mol	the amount of moles of titrant added		
Each one of this elements is a suitable xval for plotting an aquaenv object generated by				
titration.				

#### 3.4.1 Titration with HCl

The standard titration type is titration with hydrochloric acid (HCl). A simple example will illustrate this best.

An object of type aquaenv needs to be created to define the initial conditions of the titration. That is temperature, salinity, depth, the concentrations of all summed quantities and the initial pH (or [TA]).

```
> ae_init <- aquaenv(Tc = 15, S = 35, SumCO2 = 0.0035, + SumNH4 = 2e-05, pH = 11.3)
```

Then titration can be run to create the object describing the simulated titration. In this example the titrant is HCl of the relatively low concentration of 0.01 mol/kg-solution. The

sample solution amounts to 10 g. To sweep a considerable pH range quite a lot of sample needs to be added: 20 g. This means the salinity of the solution in the titration vessel will change due to dilution with the titrant solution. For this reason, the salinity of the titrant solution needs to be given via the parameter S\_titrant. However, we assume the titrant does not contain borate, sulfate or fluoride, that is why we do not set the flag seawater\_titrant to TRUE.

```
> ae <- titration(ae_init, mass_sample = 0.01, mass_titrant = 0.02,
+ conc_titrant = 0.01, S_titrant = 0.5, steps = 100)</pre>
```

To get a quick overview, all elements of the obtained aquaenv object can be plotted

```
> plot(ae, xval = ae$delta_mass_titrant, xlab = "HCl solution added [kg]", + mfrow = c(10, 10), newdevice = FALSE)
```

Then, a selection of elements can be plotted as a function of the added titrant mass,

```
> what <- c("TA", "pH", "CO2", "HCO3", "CO3", "BOH3", "BOH4",
+ "OH", "NH4", "NH3", "H2SO4", "HSO4", "SO4", "HF",
+ "F", "pCO2")
> plot(ae, xval = ae$delta_mass_titrant, xlab = "HCl solution added [kg]",
+ what = what, size = c(12, 8), mfrow = c(4, 4), newdevice = FALSE)
```

titrant concentration offset, or the moles of added titrant

```
> plot(ae, xval = ae\$delta_conc_titrant, xlab = "[HCl] offset added [mol/kg-soln]",
+ what = what, size = c(12, 8), mfrow = c(12, 8))
> plot(ae, xval = ae\$delta_moles_titrant, xlab = "HCl added [mol]",
+ what = what, size = c(12, 8), mfrow = c(12, 8))
```

However, it is also possible to plot this selection of elements against the calculated free scale pH

```
> plot(ae, xval = aepH, xlab = "free scale pH", what = what,
+ size = c(12, 8), mfrow = c(4, 4), newdevice = FALSE)
```

As mentioned earlier, the function plot.aquaenv offers the possibility of creating bjerrum plots from objects obtained with titration. The simplest way to do that is (remember the newdevice=FALSE is just needed to produce plots that are nicely woven into this vignette)

```
> plot(ae, bjerrum = TRUE, newdevice = FALSE)
```

Or just select a few elements

```
> what <- c("CO2", "HCO3", "CO3")
> plot(ae, what = what, bjerrum = TRUE, newdevice = FALSE)
```

Again, the plotting style can be customized

```
> plot(ae, what = what, bjerrum = TRUE, lwd = 4, palette = c("cyan",
+ "magenta", "yellow"), bg = "gray", legendinset = 0.1,
+ legendposition = "topleft", newdevice = FALSE)
```

However, generally Bjerrum plots are done on the log scale. This can be accomplished using the flag log

```
> what <- c("CO2", "HCO3", "CO3", "BOH3", "BOH4", "OH",
+ "NH4", "NH3", "H2SO4", "HSO4", "SO4", "HF", "F")
> plot(ae, what = what, bjerrum = TRUE, log = TRUE, newdevice = FALSE)
```

Furthermore, we can zoom in to the region of most interest to marine scientists

```
> plot(ae, what = what, bjerrum = TRUE, log = TRUE, ylim = c(-6, + -1), legendinset = 0, lwd = 3, palette = c(1, 3, + 5, 6, colors()[seq(100, 250, 6)]), newdevice = FALSE)
```

#### 3.4.2 Titration with NaOH

Similar to the titration with HCl, also a titration with NaOH can be simulated

```
> ae <- titration(aquaenv(Tc = 15, S = 35, SumCO2 = 0.0035,
+ SumNH4 = 2e-05, pH = 2), mass_sample = 0.01, mass_titrant = 0.02,
+ conc_titrant = 0.01, S_titrant = 0.5, steps = 50,
+ type = "NaOH")</pre>
```

Plottinge everything

```
> plot(ae, xval = ae$delta_mass_titrant, xlab = "NaOH solution added [kg]", + mfrow = c(10, 10))
```

Plotting selectively

```
> what <- c("TA", "pH", "CO2", "HCO3", "CO3", "BOH3", "BOH4",
+ "OH", "NH4", "NH3", "H2SO4", "HSO4", "SO4", "HF",
+ "F", "pCO2")
> plot(ae, xval = ae$delta_mass_titrant, xlab = "NaOH solution added [kg]",
+ what = what, size = c(12, 8), mfrow = c(4, 4))
> plot(ae, xval = ae$pH, xlab = "free scale pH", what = what,
+ size = c(12, 8), mfrow = c(4, 4))
```

Bjerrum plots

```
> what <- c("CO2", "HCO3", "CO3")
> plot(ae, what = what, bjerrum = TRUE, newdevice = FALSE)

> what <- c("CO2", "HCO3", "CO3", "BOH3", "BOH4", "OH",
+ "NH4", "NH3", "H2SO4", "HSO4", "SO4", "HF", "F")
> plot(ae, what = what, bjerrum = TRUE, log = TRUE, ylim = c(-6,
+ -1), legendinset = 0, lwd = 3, palette = c(1, 3,
+ 4, 5, 6, colors()[seq(100, 250, 6)]), newdevice = FALSE)
```

# 3.4.3 Titration with a titrant with high concentrations and a large sample volume - classical Bjerrum plots

The Bjerrum plots created in the previous two sections do not really look like the classical textbook ones. This is because we simulated a titration with a small sample volume and a titrant with low concentrations. As a result the total concentrations like, e.g., total carbonate decreased due to dilution. In simulating a titration with a rather large volume and a titrant with high concentrations the volume and salinity corrections do not matter any more and graphs known from textbooks (e.g. Zeebe and Wolf-Gladrow 2001) are produced.

```
> ae \leftarrow titration(aquaenv(Tc = 15, S = 35, SumCO2 = 0.0035,
      SumNH4 = 2e-05, pH = 11.3), mass_sample = 100, mass_titrant = 0.5,
      conc_titrant = 3, S_titrant = 0.5, steps = 100)
Plotting everything
> plot(ae, xval = ae$delta_mass_titrant, xlab = "HCl solution added [kg]",
      mfrow = c(10, 10))
Plotting selectively and with different elements for xval
> what <- c("TA", "pH", "CO2", "HCO3", "CO3", "BOH3", "BOH4",
      "OH", "NH4", "NH3", "H2SO4", "HSO4", "SO4", "HF",
      "F", "pCO2")
> plot(ae, xval = ae$delta_mass_titrant, xlab = "HCl solution added [kg]",
      what = what, size = c(12, 8), mfrow = c(4, 4))
> plot(ae, xval = ae$pH, xlab = "free scale pH", what = what,
      size = c(12, 8), mfrow = c(4, 4))
> plot(ae, xval = ae$delta_conc_titrant, xlab = "[HCl] offset added [mol/kg-soln]",
      what = what, size = c(12, 8), mfrow = c(4, 4))
> plot(ae, xval = ae$delta_moles_titrant, xlab = "HCl added [mol]",
      what = what, size = c(12, 8), mfrow = c(4, 4))
Creating different kinds of Bjerrum plots
> plot(ae, bjerrum = TRUE)
> what <- c("CO2", "HCO3", "CO3")</pre>
> plot(ae, what = what, bjerrum = TRUE)
> plot(ae, what = what, bjerrum = TRUE, lwd = 4, palette = c("cyan",
      "magenta", "yellow"), bg = "gray", legendinset = 0.1,
      legendposition = "topleft")
> what <- c("CO2", "HCO3", "CO3", "BOH3", "BOH4", "OH",
      "NH4", "NH3", "H2SO4", "HSO4", "SO4", "HF", "F")
> plot(ae, what = what, bjerrum = TRUE, log = TRUE)
and the classical textbook one
> plot(ae, what = what, bjerrum = TRUE, log = TRUE, ylim = c(-6),
      -1), legendinset = 0, lwd = 3, palette = c(1, 3, 3)
      4, 5, 6, colors()[seq(100, 250, 6)]), newdevice = FALSE)
```

# 3.5 Calculating information from titration curves: the function TAfit

## 3.5.1 A little theory

While titrating a sample of natural seawater with HCl there one sees two clear equivalence points (Dickson 1981) The second equivalence point is the equivalence point of total alkalinity and the difference between the second and the first equivalence point signifies the total amount of  $\sum CO_2$  of the sample Hansson and Jagner (1973).

This can be illustrated with <code>AquaEnv</code>. The respective titration curve can be plotted, together with its first and second derivative. Furthermore, the equivalence points can be marked with vertical lines (Please note that for a titrant concentration of 0.01 mol/kg-solution and 0.01 kg of sample, the value of the concentration (in mol/kg-solution) of total alkalinity and total carbonate equals the value of the total amount (in mol)).

```
> ae_init <- aquaenv(Tc = 15, S = 35, SumCO2 = 0.0035,
      SumNH4 = 2e-05, pH = 11.3)
> ae <- titration(ae_init, mass_sample = 0.01, mass_titrant = 0.02,
      conc_titrant = 0.01, S_titrant = 0.5, steps = 100)
> plot(ae, xval = ae$delta_mass_titrant, xlab = "HCl solution added [kg]",
      what = "pH", xlim = c(0, 0.015), newdevice = FALSE)
> par(new = TRUE)
> plot(ae$delta_mass_titrant[1:100], diff(ae$pH), type = "l",
      col = "red", xlim = c(0, 0.015), ylab = "", xlab = "",
      yaxt = "n")
> par(new = TRUE)
> plot(ae$delta_mass_titrant[2:100], diff(diff(ae$pH)),
      type = "1", col = "green", xlim = c(0, 0.015), ylab = "",
      xlab = "", yaxt = "n")
> abline(h = 0, col = "green")
> abline(v = ae$TA[[1]])
> abline(v = ae$TA[[1]] - ae$SumCO2[[1]])
```

Following classical chemical textbooks (e.g. Skoog and West 1982), one can determine [TA] and  $[\sum CO_2]$  of a sample by graphically determining those equivalence points. However, there is no mechanistic understanding of the contents of the solution involved in doing so.

Other methods, called "Gran evaluations" (Gran 1952; Hansson and Jagner 1973; Dickson 1981; Haraldsson, Anderson, Hassellov, Hulth, and Olsson 1997; Anderson, Turner, Wedborg, and Dyrssen 1999), try to linearize the mechanistic model of what is going on in the solution during titration. They define the so called linear "Gran functions" and try to find their roots to determine the equivalence points. We will illustrate that by plotting the Gran functions F0 (blue) and F2 (-F1, green) and again mark the equivalence points with vertical lines. The y=zero line for the Gran functions is indicated by a horizontal line

```
> plot(ae, xval = ae$delta_mass_titrant, xlab = "HCl solution added [kg]",  
+ what = "pH", xlim = c(0, 0.015), newdevice = FALSE)
```

```
> prot1 <- c()
> for (i in 1:length(ae$pH)) {
     prot1 <- c(prot1, (10^-(ae$pH[[i]]) + ae$HSO4[[i]] +</pre>
          ae$HF[[i]] + ae$CO2[[i]] - ae$CO3[[i]] - ae$BOH4[[i]] -
          ae$OH[[i]]))
+ }
> par(new = TRUE)
> plot(ae$delta_mass_titrant, prot1, type = "1", col = "blue",
      xlim = c(0, 0.015), ylab = "", xlab = "", yaxt = "n",
      ylim = c(-0.015, 0.015))
> prot2 <- c()
> for (i in 1:length(ae$pH)) {
     prot2 <- c(prot2, (10^-(ae$pH[[i]]) + ae$HSO4[[i]] +</pre>
          ae$HF[[i]] - ae$HCO3[[i]] - 2 * ae$CO3[[i]] -
          ae$BOH4[[i]] - ae$OH[[i]]))
+ }
> par(new = TRUE)
> plot(ae$delta_mass_titrant, prot2, type = "l", col = "green",
      xlim = c(0, 0.015), ylab = "", xlab = "", yaxt = "n",
      ylim = c(-0.015, 0.015))
> abline(v = ae$TA[[1]])
> abline(v = ae$TA[[1]] - ae$SumCO2[[1]])
> abline(h = 0)
```

One can see that the Gran functions actually are not linear. This is due to volume and salinity change effects during the titration. This can be overcome by either employing "modified Gran functions" (see Haraldsson *et al.* 1997) that correct for the volume changes or by using a titration with a titrant with high concentrations and a large sample volume (Please not that here the value of the concentration of total alkalinity and total carbonate does not equal their total amount)

```
> ae <- titration(aquaenv(Tc = 15, S = 35, SumCO2 = 0.0035,
      SumNH4 = 2e-05, pH = 11.3), mass_sample = 100, mass_titrant = 0.5,
      conc_titrant = 3, S_titrant = 0.5, steps = 100)
> plot(ae, xval = ae$delta_mass_titrant, xlab = "HCl solution added [kg]",
      what = "pH", xlim = c(0, 0.5), newdevice = FALSE)
> prot1 <- c()
> for (i in 1:length(ae$pH)) {
     prot1 <- c(prot1, (10^-(ae$pH[[i]]) + ae$HSO4[[i]] +</pre>
          ae$HF[[i]] + ae$CO2[[i]] - ae$CO3[[i]] - ae$BOH4[[i]] -
          ae$OH[[i]]))
+ }
> par(new = TRUE)
> plot(ae$delta_mass_titrant, prot1, type = "l", col = "blue",
      xlim = c(0, 0.5), ylab = "", xlab = "", yaxt = "n",
     ylim = c(-0.015, 0.015))
> prot2 <- c()
```

Another proposed method of determining [TA] and  $[\sum CO_2]$  is to not only determine the two equivalence points, but to fit the whole titration curve with a theoretical titration curve based on a mechanistic model of what is going on in the solution during the titration (Dickson 1981; DOE 1994; Anderson *et al.* 1999). The function titration of **AquaEnv** provides exactly such a theoretical titration curve and the function Tafit makes use of this fact to determine [TA] and  $[\sum CO_2]$  of a sample by non linear curve fitting.

# 3.5.2 Determining [TA] and $[\sum CO_2]$ by non linear curve fitting

**3.5.2.1 Proof of concept** First, a proof of concept will show that the function TAfit is implemented consistently. Some "data" can be generated with the titration function.

```
> initial_ae <- aquaenv(Tc = 15, S = 35, SumCO2 = 0.002,
+ TA = 0.0022)
> ae <- titration(initial_ae, mass_sample = 0.01, mass_titrant = 0.003,
+ conc_titrant = 0.01, S_titrant = 0.5, steps = 20)
```

Now, the input data for the  $\mathtt{TAfit}$  routine can be generated: a table with the added mass of the titrant and the resulting free scale  $\mathtt{pH}$ 

```
> titcurve <- cbind(ae$delta_mass_titrant, ae$pH)</pre>
```

Note that For the TAfit all total quantities except SumCO2 (SumNH4, SumH2S, SumH3PO4, SumSiOH4, SumHNO3, SumHNO2, SumBOH3, SumH2SO4, SumHF) need to be known. However, the latter three can be calculated from salinity as it is done in this example.

```
> fit1 <- TAfit(initial_ae, titcurve, conc_titrant = 0.01,
+ mass_sample = 0.01, S_titrant = 0.5)
> fit1
```

Thus, we see that TAfit calculates the correct SumCO2 and TA values.

TAfit can also take E (V) values as input variables, so we generate E values using E0=0.4 V and the Nernst equation. However, to do so we first need to convert our pH curve to the

seawater pH scale. According to (DOE 1994, p.7, ch.4, sop.3), the Nernst equation relates E to the total proton concentration, but, if fluoride is present, its effect (as proton donor/acceptor) is also measured. Hence, we use the seawater scale here.

```
> swstitcurve <- convert(titcurve[, 2], "pHscale", "free2sws",
+ Tc = Tc, S = S)
> Etitcurve <- cbind(titcurve[, 1], (0.4 - ((Constants$R/10) *
+ initial_ae$Tk/Constants$F) * log(10^-swstitcurve)))</pre>
```

Again, TAfit can be executed, this time also calculating E0. Note that the flag verbose=TRUE causes TAfit to show the progress of the fitting procedure in a plot window.

```
> fit2 <- TAfit(initial_ae, Etitcurve, conc_titrant = 0.01,
+ mass_sample = 0.01, S_titrant = 0.5, Evals = TRUE,
+ verbose = TRUE)
> fit2
```

Furthermore, TAfit can fit K\_CO2 as well, however, one single value for the whole titration curve is fitted, i.e. there is no correction for K\_CO2 changes due to changing S due to mixing with the titrant

```
> fit3 <- TAfit(initial_ae, titcurve, conc_titrant = 0.01,
+ mass_sample = 0.01, S_titrant = 0.5, K_CO2fit = TRUE)
> fit3
> initial_ae$K_CO2
```

One can see that the fitted value for K\_CO2 is not the same as the value in the initial aquaenv object, which is the "correct" value. That is, because during data creation K\_CO2 changed along the course of the titration due to changes in salinity. Assuming that the titrant has the same salinity as the sample (and is made up of natural seawater, i.e. containing SumBOH4, SumH2SO4 and SumHF as functions of S), then the "correct" K\_CO2 should be fitted. This can be accomplished in TAfit by not giving the argument S\_titrant (i.e. assuming the titrant has the same salinity as the sample) and setting the flag seawater\_titrant to TRUE

```
> ae <- titration(initial_ae, mass_sample = 0.01, mass_titrant = 0.003,
+ conc_titrant = 0.01, steps = 20, seawater_titrant = TRUE)
> titcurve <- cbind(ae$delta_mass_titrant, ae$pH)
> fit4 <- TAfit(initial_ae, titcurve, conc_titrant = 0.01,
+ mass_sample = 0.01, K_CO2fit = TRUE, seawater_titrant = TRUE)
> fit4
```

Furthermore, TA, SumCO2, K\_CO2 and E0 can be fitted at the same time.

```
> Etitcurve <- cbind(titcurve[, 1], (0.4 - ((Constants$R/10) *
+     initial_ae$Tk/Constants$F) * log(10^-titcurve[, 2])))
> fit5 <- TAfit(initial_ae, Etitcurve, conc_titrant = 0.01,
+     mass_sample = 0.01, K_CO2fit = TRUE, seawater_titrant = TRUE,
+     Evals = TRUE)
> fit5
```

Sometimes, the obtained titration curve is not equally spaced on the x axis. TAfit can deal with such curves if the flag equalspaced is set to FALSE

```
> neqsptitcurve <- rbind(titcurve[1:9, ], titcurve[11:20,
+     ])
> fit6 <- TAfit(initial_ae, neqsptitcurve, conc_titrant = 0.01,
+     mass_sample = 0.01, seawater_titrant = TRUE, equalspaced = FALSE,
+     verbose = TRUE, debug = TRUE)
> fit6

Finally, some "noise" is added to the generated data
```

```
> noisetitcurve <- titcurve * rnorm(length(titcurve), mean = 1,
+ sd = 0.01)
> fit7 <- TAfit(initial_ae, noisetitcurve, conc_titrant = 0.01,
+ mass_sample = 0.01, seawater_titrant = TRUE, verbose = TRUE)
> fit7
```

The flag verbose=TRUE prompts to show the traject of the fitting procedure in a plot window. However, each new fit is plotted over the first one and Sweave includes only the first plot in each code chunk in the resulting LATEX file. Therefore, we use the flag debug=TRUE to visualize the final fit

```
> ylim = range(noisetitcurve[, 2], calc)
> xlim = range(tit$delta_mass_titrant, noisetitcurve[,
+ 1])
> plot(noisetitcurve[, 1], noisetitcurve[, 2], xlim = xlim,
+ ylim = ylim, type = "l", xlab = "delta mass titrant",
+ ylab = "pH (free scale)")
> par(new = TRUE)
> plot(tit$delta_mass_titrant, calc, xlim = xlim, ylim = ylim,
+ type = "l", col = "red", xlab = "", ylab = "")
```

**3.5.2.2 Test with generated data from literature** Dickson (1981) provided a synthetic dataset to test total alkalinity fitting programs. This dataset is included in **AquaEnv** as sample\_dickson. Following quantities are given

```
> conc_titrant <- 0.3
> mass_sample <- 0.2
> S_titrant <- 14.835
> SumBOH3 <- 0.00042
> SumH2SO4 <- 0.02824
> SumHF <- 7e-05</pre>
```

Note that all concentrations are in mol/kg-solution and the mass of the sample is in kg. Note further that the salinity of the titrant has been calculated from its ionic strength of 0.3

mol/kg-soln.

In the original dataset as represented in sample\_dickson, the mass of titrant is given in g which needs to be converted to kg

```
> sam <- cbind(sample_dickson[, 1]/1000, sample_dickson[,
+ 2])</pre>
```

Then an attempt to recalculate the [TA] and  $[\sum CO_2]$  values given in Dickson (1981) ([TA]=0.00245 mol/kg-soln and  $[\sum CO_2]$  0.00220 mol/kg-soln) can be done

```
> dicksonfit <- TAfit(aquaenv(Tc = 25, S = 35, SumBOH3 = SumBOH3,
+ SumH2SO4 = SumH2SO4, SumHF = SumHF), sam, conc_titrant,
+ mass_sample, S_titrant = S_titrant, debug = TRUE)
> dicksonfit

$TA
[1] 0.002464256
attr(,"unit")
[1] "mol/kg-soln"
```

# \$SumCO2

[1] 0.002191958
attr(,"unit")
[1] "mol/kg-soln"

#### \$sumofsquares

[1] 0.01287255

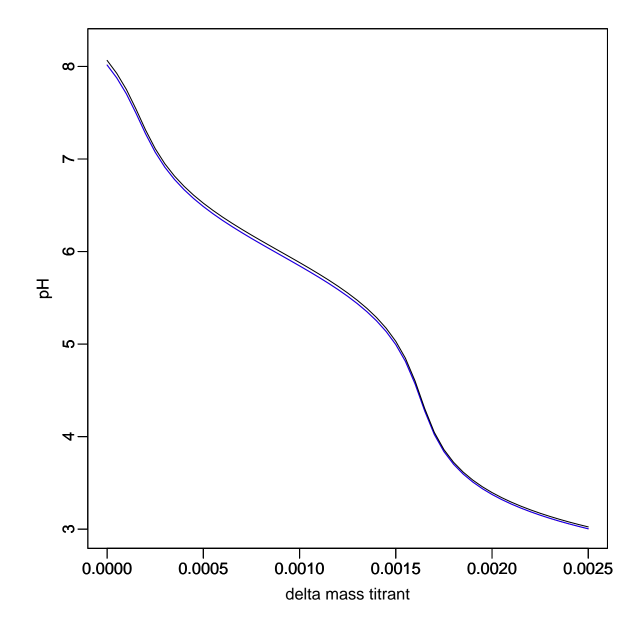
This shows the fit is not accurate. Why is that so?

#### 3.5.2.2.1 Does the salinity correction (S\_titrant) matter?

Let us calculate a theoretical titration without salinity correction

Now the difference between both curves (in red and blue) and the "Dickson" curve (in black) can be visualized

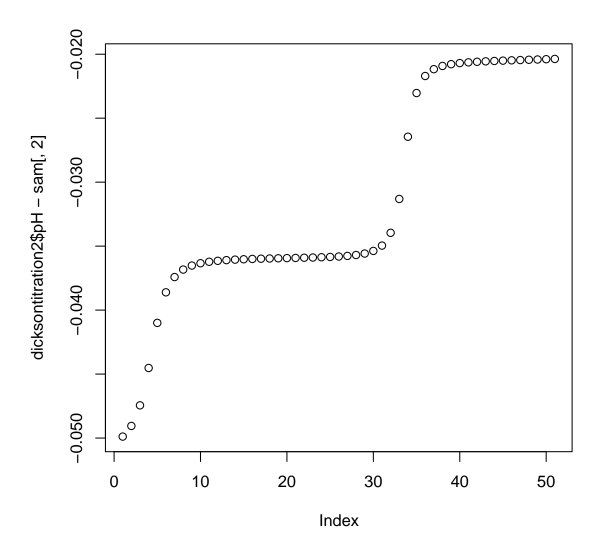
```
> plot(dicksontitration1, xval = dicksontitration1$delta_mass_titrant,
+ what = "pH", xlim = c(0, 0.0025), ylim = c(3, 8.2),
+ newdevice = FALSE, col = "red", xlab = "delta mass titrant")
> par(new = TRUE)
> plot(dicksontitration2, xval = dicksontitration2$delta_mass_titrant,
+ what = "pH", xlim = c(0, 0.0025), ylim = c(3, 8.2),
+ newdevice = FALSE, col = "blue", xlab = "")
> par(new = TRUE)
> plot(sam[, 1], sam[, 2], type = "l", xlim = c(0, 0.0025),
+ ylim = c(3, 8.2), xlab = "", ylab = "")
```



That means, the salinity correction makes no significant difference (the red and the blue curve cannot be discerned), because the relation between the total amount of sample and the added amount of titrant is very large: salinity only drops from 35 to 34.75105.

But there is an offset between the "Dickson" curve and our curve

> plot(dicksontitration2\$pH - sam[, 2])



# 3.5.2.2.2 Does fitting K\_CO2 as well improve the fit?

```
> dicksonfit2 <- TAfit(aquaenv(Tc = 25, S = 35, SumBOH3 = SumBOH3,
```

- + SumH2SO4 = SumH2SO4, SumHF = SumHF), sam, conc\_titrant,
- + mass\_sample, S\_titrant = S\_titrant, debug = TRUE,

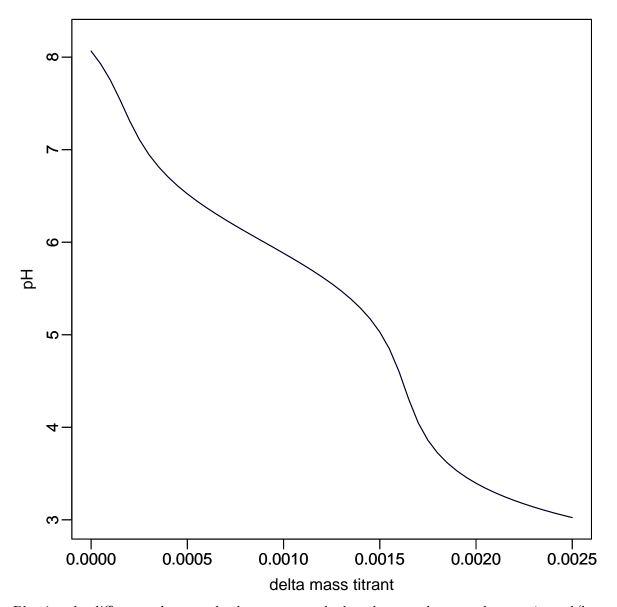
```
K_{CO2fit} = TRUE
> dicksonfit2
$TA
[1] 0.002458081
attr(,"unit")
[1] "mol/kg-soln"
$SumCO2
[1] 0.002194006
attr(,"unit")
[1] "mol/kg-soln"
$K_C02
[1] 1.030960e-06
attr(,"unit")
[1] "mol/kg-soln"
attr(,"pH scale")
[1] "free"
$sumofsquares
[1] 0.005724457
```

Yes it does, but it is not optimal yet.

There still remains one major difference between the calculations carried out in Dickson (1981) and the calculations in **AquaEnv**: Dickson (1981) uses fixed values for the equilibrium constants and does not calculate them as functions of temperature and salinity. Furthermore, the values that are used in Dickson (1981) are not exactly the same as are obtained in **AquaEnv** for the same salinity and temperature.

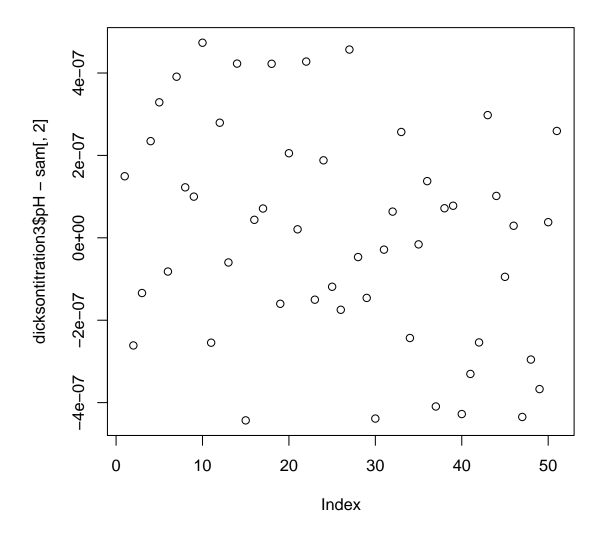
Let us calculate a theoretical titration curve employing exactly the same equilibrium constant values as used in Dickson (1981) and plot the result together with the "Dickson" curve

```
> par(new = TRUE)
> plot(sam[, 1], sam[, 2], type = "1", xlim = c(0, 0.0025),
+ ylim = c(3, 8.2), xlab = "", ylab = "")
```



Plotting the differences between both curves reveals that they are the same down to 1 umol/kg-soln.

> plot(dicksontitration3\$pH - sam[, 2])



Calculating [TA] and  $[\sum CO_2]$  using TAfit and exactly the same equilibrium constant values as used in Dickson (1981)

```
> dicksonfit3 <- TAfit(aquaenv(Tc = 25, S = 35, SumBOH3 = SumBOH3,
+ SumH2SO4 = SumH2SO4, SumHF = SumHF, k_w = 4.32e-14,
+ k_co2 = 1e-06, k_hco3 = 8.2e-10, k_boh3 = 1.78e-09,
+ k_hso4 = (1/12.3), k_hf = (1/408)), sam, conc_titrant,
+ mass_sample, S_titrant = S_titrant, debug = TRUE,
+ k_w = 4.32e-14, k_co2 = 1e-06, k_hco3 = 8.2e-10,
+ k_boh3 = 1.78e-09, k_hso4 = (1/12.3), k_hf = (1/408))
> dicksonfit3
$TA
[1] 0.00245
```

```
attr(,"unit")
[1] "mol/kg-soln"

$SumCO2
[1] 0.0022
attr(,"unit")
[1] "mol/kg-soln"

$sumofsquares
[1] 3.280551e-12
```

reveals that now exactly the same values are calculated as are given in Dickson (1981).

# A References for the elements of an object of class aquaenv

element	references
Cl	(DOE 1994, chapter 5, p. 11), and (Zeebe and Wolf-Gladrow 2001, p. 100, footnote 3)
I	(DOE 1994, chapter 5, p. 13, 15), (Zeebe and Wolf-Gladrow 2001, p.12), and (Roy et al. 1993b, p.257). Note that the approximation $I/(mol/kg\text{-solution}) \approx 0.0199201$ S is given in (Millero 1982, p. 428.). This relationship converted into mol/kg-H <sub>2</sub> O and the last digits adjusted (from 0.0199201 to 0.019924) results in the formula used here.
density	Millero and Poisson (1981) and (DOE 1994, chapter 5, p. 6f).
SumBr, ClConc, Na, Mg, Ca, K, Sr	(DOE 1994, chapter 5, p.11)
molal2molin	(Roy et al. 1993b, p.257), and (DOE 1994, chapter 5, p. 15)
$\begin{array}{c} {\rm free2tot}, \\ {\rm tot2free} \end{array}$	(Dickson 1984, p.2302), (DOE 1994, chapter 4, p.16), (Zeebe and Wolf-Gladrow 2001, p.57, 261)
free2sws, tot2sws, sws2free, sws2tot	(Dickson 1984, p.2303), (Zeebe and Wolf-Gladrow 2001, p.57)
K0_CO2	Weiss (1974), (DOE 1994, chapter 5, p. 13) (here it is stated that the unit is mol/(kg-solution*atm)), (Millero 1995, p.663), (Zeebe and Wolf-Gladrow 2001, p.257)
K0_O2	derived from a formula for the oxygen saturation concentration in ml-O <sub>2</sub> /kg-solution by Weiss (1970) using the first virial coefficient of oxygen (Atkins 1996, p. 41, 1029) and the atmospheric oxygen fugacity (Williams 2004)
K_W	(Millero 1995, p.670) ( <b>original reference</b> , but slightly different formula for seawater pH), (DOE 1994, chapter 5, p. 18) (NOT the original reference! DOE (1994) cites in an update from 1997 Millero (1995)! However the version of the formula used here is the one converted to total pH scale given in DOE (1994)), and (Zeebe and Wolf-Gladrow 2001, p. 258). Constant type (stoichiometric), pH scale (total, converted to free here), and concentration unit (mol/kg-solution squared): (DOE 1994, chapter 5, p. 12,18), pH scale also in (Zeebe and Wolf-Gladrow 2001, p. 258).
K_HSO4	(DOE 1994, chapter 5 page 13), (Zeebe and Wolf-Gladrow 2001, p. 260), Dickson (1990b) (original reference). Constant type (stoichiometric), pH scale (free), and concentration unit (mol/kg-H <sub>2</sub> O converted to mol/kg-solution here): (DOE 1994, chapter 5, p. 13).

 $K_{-}HF$ 

(Dickson and Riley 1979b, p. 91) (original reference), (DOE 1994, c. 5, p. 15), (Roy et al. 1993b, p. 257), (Dickson and Millero 1987, p. 1783), (Millero 1995, p. 664), (Zeebe and Wolf-Gladrow 2001, p. 260) (converted to molinty and total scale). Constant type (stoichiometric), pH scale (free), and concentration unit (mol/kg-H<sub>2</sub>O converted to mol/kg-solution here): (DOE 1994, chapter 5, p. 15, 16).

K\_CO2. K\_HCO3 (Roy et al. 1993b, p. 254) (original reference), (DOE 1994, chapter 5, p.14) (in a version converted to mol/kg-H<sub>2</sub>O), (Millero 1995, p. 664), (Zeebe and Wolf-Gladrow 2001, p. 255). Constant type (stoichiometric) and concentration unit (mol/kg-H<sub>2</sub>O converted to mol/kg-solution here): (DOE 1994, chapter 5, p. 14, 15), pH scale (total, converted to free here): In (DOE 1994, chapter 5, p. 12) the total scale is stated for the formula for high salinities and thus can be inferred for the formula for low salinities. The scale is also indirectly stated for both formulations in the original reference Roy et al. (1993b). Note that in Roy et al. (1993b) a function for pure water (Millero 1979) is cited and a function for seawater is derived. In Millero (1995) it is stated that for S<5 the fresh water formula of (Millero 1979) should be used and for S>=5 the seawater formula derived in Roy et al. (1993b). However, both formulations do not always intersect at S=5. The true intersection with respect to salinity S is a function of temperature Tk. Here, we first calculate this intersection by numerical root finding and then decide which formulation to use. This practise results in a continuous function with respect to S. (Note that Millero (1979) is restated wrongly in Roy et al. (1993b): one of the numerical values for the function for  $K_{CO_2}^*$  is given as 310.48919, but correct is 2310.48919. However, in Millero (1995) this value is stated correctly.)

K\_BOH3

(Dickson 1990a, p. 763) (original, but mol/kg-H<sub>2</sub>O version), (DOE 1994, ch. 5, p. 14), (Zeebe and Wolf-Gladrow 2001, p. 262), (Millero 1995, p.669) (mol/kg-H<sub>2</sub>O version), agrees with data in Roy, Roy, Lawson, Vogel, Moore, Davis, and Millero (1993a). Constant type (stoichiometric) and concentration unit (mol/kg-solution): (DOE 1994, chapter 5, p. 14), pH scale (total): (DOE 1994, chapter 5, p. 12) and (Zeebe and Wolf-Gladrow 2001, p.263).

 $K_NH4$ 

Millero et al. (1995) (original reference), (Millero 1995, p.671). Constant type (stoichiometric) and concentration unit (mol/kg-solution): (Millero 1995, p.671), pH scale (seawater, converted to free here): Lewis and Wallace (1998) (in corrections of Millero (1995)).

 $K_{-}H2S$ 

Millero et al. (1988) (original reference), (Millero 1995, p.671). Constant type (stoichiometric) and concentration unit (mol/kg-solution): (Millero 1995, p.671), pH scale (seawater, converted to free here): Lewis and Wallace (1998) (in corrections of Millero (1995)).

K\_H3PO4, K\_H2PO4, K\_HPO4

(Millero 1995, p.670) (original reference, but formula for seawater scale pH), (DOE 1994, ch. 5, p 16,17), agrees with data in Dickson and Riley (1979a). Constant type (stoichiometric), concentration unit (mol/kg-solution), and pH scale (total, converted to free here): (DOE 1994, chapter 5, p. 12, 16, 17).

 $K_SiOH4$ 

Millero et al. (1988) (original reference), (DOE 1994, chapter 5, p 17), (Millero 1995, p.671) (formula for seawater scale pH) Constant type (stoichiometric), concentration unit (mol/kg-H<sub>2</sub>O converted to mol/kg-solution here by omitting the conversion summand ln(1-0.001005 S)), and pH scale (total, converted to free here): (DOE 1994, chapter 5, p. 12, 17).

K\_SiOOH3

Wischmeyer et al. (2003) (original reference), corrected due to personal communication with Dieter Wolf-Gladrow (one of the authors). The corrected version can be obtained from either Dieter Wolf-Gladrow or Andreas F Hofmann (a.hofmann@nioo.knaw.nl). Constant type (stoichiometric), concentration unit (mol/kg-solution), and pH scale (total, converted to free here): Wischmeyer et al. (2003).

 $K_{-}HNO2$ 

Constant value, not a function of temperature and salinity! Obtained as a hybrid pk value (featuring the activity of the proton but the concentration of other species (see Zeebe and Wolf-Gladrow (2001) for a treatment of different types of equilibrium constants) in molar concentration (mol/l) on the NBS pH scale (Durst 1975) from Riordan et al. (2005). Used as an approximation for the stoichiometric  $K_{HNO_2}^*$  in mol/kg-solution on the free proton pH scale here.

Constant value, not a function of temperature and salinity! Obtained as a standard pK value from (Atkins 1996, p. 1045). Used as an approximation for the stoichiometric  $K_{12SO_4}^*$  in mol/kg-solution on the free proton pH scale here.

 $K_{-}HS$ 

Constant value, not a function of temperature and salinity! Obtained as a standard pK value from (Atkins 1996, p. 1045). Used as an approximation for the stoichiometric  $K_{Hs^-}^*$  in mol/kg-solution on the free proton pH scale here.

 $K_H2SO4$ 

Mucci (1983) (original reference), Boudreau (1996). Note that in there are errors in Boudreau Ksp\_calcite, (1996):  $b_0$  for calcite is not 0.7712 but 0.77712 and  $b_1$  for arganite is not 0.001727 but 0.0017276. Ksp\_aragonite As given in Dickson (1984), p. 2303 (use of "m") and Dickson and Riley (1979a), p. 91f all Нq concentrations appearing in the definition of the total and the seawater pH scale are molal (mol/kg-H<sub>2</sub>O) concentrations. But in Roy et al. (1993b), p. 257 and in DOE (1994), chapter 4, SOP 6, p. 1 it is stated, that concentrations for the seawater and total pH scale are in mol/kg-solution. To be consistent with DOE (1994) molin concentrations (mol/kg-solution) are chosen for calculating the revelle (Zeebe and Wolf-Gladrow 2001, p.73) dTAdKdKdS, Hofmann et al. (2008a) dTAdKdKdT, dTAdKdKdd, dTAdKdKd-SumH2SO4, dTAdKdKd-SumHF

The values for K\_W, K\_HSO4, K\_HF, K\_CO2, K\_HCO3, K\_BOH3, K\_NH4, K\_H2S, K\_H3PO4, K\_H2PO4, K\_HPO4, K\_SiOH4, K\_SiOOH3, Ksp\_calcite, Ksp\_aragonite obtained as functions of salinity S and temperature Tc from the above references are pressure corrected using the given depth d and the calculated hydrostatic pressure hydroP according to Millero (1995) with corrections by Lewis and Wallace (1998).

In general it is to be said that all corrections from Lewis and Wallace (1998) have been applied.

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