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Improving noble gas based paleoclimate reconstruction and groundwater dating using ²⁰Ne/²²Ne ratios

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Abstract—The interpretation of noble gas concentrations in groundwater with respect to recharge temperature and fractionated excess gas leads to different results on paleo-climatic conditions and on residence times depending on the choice of the gas partitioning model. Two fractionation models for the gas excess are in use, one assuming partial re-equilibration of groundwater supersaturated by excess air (PR-model, Stute et al., 1995), the other assuming closed-system equilibration of groundwater with entrapped air (CE-model, Aeschbach-Hertig et al., 2000). In the example of the Continental Terminal aquifers in Niger, PR- and CE- model are both consistent with the data on elemental noble gas concentrations (Ne, Ar, Kr, and Xe). Only by including the isotope ratio ${}^{20}\text{Ne}/{}^{22}\text{Ne}$ it can be demonstrated that the PR-model has to be rejected and the CE-model should be applied to the data. In dating applications ³He of atmospheric origin (³He_{atm}) required to calculate ³H-³He water ages is commonly estimated from the Ne excess presuming that gas excess is unfractionated air (UA-model). Including in addition to the Ne concentration the ${}^{20}\text{Ne}/{}^{22}\text{Ne}$ ratio and the concentration of Ar enables a rigorous distinction between PR-, CE- and UA-model and a reliable determination of ³He_{atm} and of ³H-³He water ages. *Copyright* © 2002 Elsevier Science Ltd

1. INTRODUCTION

Noble gas concentrations in groundwater have been employed to provide information on paleotemperatures (e.g., Aeschbach-Hertig et al., 2000; Andrews and Lee, 1979; Beyerle et al., 1998; Mazor, 1972; Stute et al., 1995; Weyhenmeyer et al., 2000) and on residence times in aquifers (³H-³He dating e.g., Schlosser et al., 1988; Schlosser et al., 1989; Tolstikhin and Kamenskiy, 1969; ⁴He dating e.g., Andrews and Lee, 1979; Solomon, 2000). In addition to applications in paleo-climate and dating studies, noble gases can serve as excellent tools to study gas exchange between the atmosphere and groundwater because noble gases are not affected by biogeochemical transformations. The interpretation of noble gas concentrations in groundwater has been improved in the last years and rigorous least squares methods are now available (Aeschbach-Hertig et al., 1999; Ballentine and Hall, 1999) which allow the estimation of e.g., recharge temperature by considering the noble gases Ne, Ar, Kr, and Xe simultaneously.

In most aquifers noble gas concentrations are higher than the concentrations expected at atmospheric solubility equilibrium. This gas excess, commonly called "excess air" (Heaton and Vogel, 1981), must be considered in the calculation of recharge temperature (Stute and Schlosser, 1993) and in the calculation of tritiogenic ³He for dating purposes (Schlosser et al., 1989). The name "excess air" however is misleading because in many aquifers (e.g., Aeschbach-Hertig et al., 2000; Stute et al., 1995) the gas excess does not have the same composition as atmospheric air but the heavier noble gases are enriched compared

to the lighter ones. In several aquifers the amount and composition of the gas excess varies systematically over time and might provide additional information on past climatic conditions related to changes in recharge dynamics (Aeschbach-Hertig et al., 2001; Stute and Talma, 1998, Beyerle et al., in press).

Although gas excess appears to be very common in groundwaters its formation is still not well understood in detail. Two different simplified models describing the formation of excess air and its fractionation (Aeschbach-Hertig et al., 2000; Stute et al., 1995) have been employed to account for the excess of noble gases in paleo-climatic studies and in ${}^{3}\text{H}{-}^{3}\text{He}$ dating applications. The two gas exchange models differ in their assumptions on how excess air is formed and on the mechanisms responsible for fractionation.

Because the choice of the gas exchange model affects the results on paleotemperatures and on ³H-³He water ages, it is necessary to decide on the basis of the available data which of the two models should be applied. However, in most ground-waters the number of elemental noble gas concentrations which can be employed in the inverse fitting procedure to simultaneously estimate recharge temperature, gas excess and its fractionation is limited to four, namely Ne, Ar, Kr and Xe. Unfortunately He, which is the most sensitive of all noble gases to the choice of the gas exchange model, usually cannot be included in the fitting procedure because it has significant additional non-atmospheric sources which cannot be quantified independently.

In this study we demonstrate that the empirical constraints on the gas exchange models can be improved significantly by considering the concentrations of noble gas isotopes and isotope ratios in addition to the commonly used elemental con-

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Table 1a. Partial pressure in the atmosphere, equilibirum concentration C^* and molecular diffusivities D of noble gases and their isotopes at different temperatures. Equilibrium concentrations are given for freshwater with S = 0 % at P = 1 atm of moist air. C^* are calculated from the solubilities for ³He, ⁴He, Ne, Ar, and Kr (Benson and Krause, 1980; Weiss, 1970; Weiss, 1971; Weiss and Kyser, 1978), and the solubilities for Xe (Clever, 1979) using the correction for moist air (Aeschbach-Hertig et al., 1999). C^* of 22 Ne and 36 Ar are calculated from the equilibrium concentration of Ne and Ar using the fractionation factors given by Beyerle et al. (2000a). Solubilities of 20 Ne and 40 Ar are assumed to correspond to the solubilities of Ne and Ar, respectively. Volume fractions in dry air are taken from Ozima and Podosek (1983) and molecular diffusivities were calculated from the empirical relations of Jähne et al. (1987). Diffusivities of Ar, 36 Ar, 40 Ar, 22 Ne and 3 He were calculated from the other noble gas diffusivites assuming that the D is inversely proportional to the square root of the atomic mass.

		C^* at 5°C	<i>C</i> * at 10°C	C^* at 20°C	<i>C</i> * at 30°C	D at 5°C	D at $10^{\circ}\mathrm{C}$	D at 20°C	<i>D</i> at 30°C	
	volume fraction	$(cm^3 \text{ STP } g^{-1})$				$(10^{-9} \text{ m}^2 \text{s}^{-1})$				
He. ⁴ He	$5.24 \cdot 10^{-6}$	$4.760 \cdot 10^{-8}$	$4.645 \cdot 10^{-8}$	$4.477 \cdot 10^{-8}$	$4.357 \cdot 10^{-8}$	5.195	5.680	6.730	7.884	
Ne	$1.818 \cdot 10^{-5}$	$2.123 \cdot 10^{-7}$	$2.017 \cdot 10^{-7}$	$1.851 \cdot 10^{-7}$	$1.724 \cdot 10^{-7}$	2.627	2.942	3.648	4.459	
Ar	$9.34 \cdot 10^{-3}$	$4.359 \cdot 10^{-4}$	$3.861 \cdot 10^{-4}$	$3.119 \cdot 10^{-4}$	$2.599 \cdot 10^{-4}$	1.646	1.870	2.381	2.984	
Kr	$1.14 \cdot 10^{-6}$	$1.057 \cdot 10^{-7}$	$9.103 \cdot 10^{-8}$	$6.967 \cdot 10^{-8}$	$5.523 \cdot 10^{-8}$	1.029	1.200	1.608	2.114	
Xe	$8.7 \cdot 10^{-8}$	$1.581 \cdot 10^{-8}$	$1.318 \cdot 10^{-8}$	$9.517 \cdot 10^{-9}$	$7.178 \cdot 10^{-9}$	0.788	0.929	1.271	1.702	
²⁰ Ne	$1.645 \cdot 10^{-5}$	$1.921 \cdot 10^{-7}$	$1.826 \cdot 10^{-7}$	$1.675 \cdot 10^{-7}$	$1.560 \cdot 10^{-7}$	2.627	2.942	3.648	4.459	
⁴⁰ Ar	$9.303 \cdot 10^{-3}$	$4.342 \cdot 10^{-4}$	$3.846 \cdot 10^{-4}$	$3.107 \cdot 10^{-4}$	$2.589 \cdot 10^{-4}$	1.646	1.870	2.381	2.984	

centrations of noble gases. In particular, including Ne isotopes proves to be very useful in distinguishing between different gas exchange models. The interpretation of data on noble gas isotopes with respect to excess air and its fractionation is demonstrated using noble gas data from the Continental Terminal aquifers in Niger (Beyerle et al., 2002).

2. THEORY AND METHODS

2.1. Gas Exchange Models and their Sensitivity to Noble Gas Concentrations

The most simple explanation for the excess of dissolved atmospheric gases in groundwater is that initially the gasconcentrations in the groundwater are in equilibrium with the atmosphere and that gas bubbles with atmospheric gas composition are trapped and completely dissolved, introducing excess gases in the same ratio as in the atmosphere into the water. This assumption, which we refer to as the unfractionated excess air (UA) model, has been used in most paleoclimatic studies based on noble gas concentrations from groundwaters (e.g., Andrews and Lee, 1979; Heaton and Vogel, 1981; Stute and Schlosser, 1993). However, recent studies have shown that the gas excess can be fractionated compared to the atmospheric gas composition. Two models have been developed to describe this situation, the partial re-equilibration model (PR-model) by Stute et al. (1995), and the closed system equilibration model (CEmodel) by Aeschbach-Hertig et al. (2000). In the following we briefly describe the underlying model assumptions and their consequences on the noble gas concentrations and fractionation.

The PR-model assumes that initially the trapped air bubbles dissolve completely, as in the UA-model. The initial excess air suggested by the model can correspond to gas concentrations which are several times larger than the atmospheric equilibrium concentrations (e.g., Stute et al., 1995). This also implies that e.g., initial oxygen levels in groundwater might be significantly above atmospheric equilibrium which would have consequences on groundwater ecology and water quality. According to the model, the gas excess in the water leads to a diffusion of gas out of the water across the groundwater table and diffusion is assumed to be at molecular level. Because the molecular diffusivities of the noble gases decrease with atomic mass, the light noble gases are lost much faster than heavier ones (Table 1a). This process increases the ratio of heavy to light noble gas concentrations in the water and thus leads to the typical fractionation pattern of the noble gas composition in the gas excess of groundwater. Some of the fractionated gas excess remains in the groundwater because infiltration and groundwater flow eventually move the water away from the air/water interface, preventing further gas loss by diffusion.

Conceptually, the model assumes a partial re-equilibration of water which was initially over-saturated by air with atmospheric composition. The effect of the PR-model on the dissolved gas concentration i can be described by (Aeschbach-Hertig et al., 1999, reformulation of the model by Stute et al., 1995):

$$C_{i}(T,S,P,A_{pr},F_{pr}) = C^{*}_{i}(T,S,P) + A_{pr} \cdot z_{i} \cdot \exp\left(-F_{pr}\frac{D_{i}}{D_{Ne}}\right) \quad (1)$$

where C_{i}^{*} is the equilibrium concentration between atmosphere and water at atmospheric pressure P, water temperature T and salinity S during air water partitioning. The equilibrium concentration is determined using the Henry coefficient of the gas considered. Details on the calculation of C_i^* are given in Aeschbach-Hertig et al. (1999). z_i is the atmospheric volume fraction of noble gas i in dry air, D_i is its molecular diffusivity, and D_{Ne} is the molecular diffusivity of Ne. A_{pr} is the amount of initial excess air and F_{pr} is the fractionation parameter which must be positve. F_{pr} can be interpreted as being proportional to the time during which the initial gas excess is lost to the atmosphere. In most cases T, A_{pr} and F_{pr} are treated as fit parameters whereas P and S are commonly prescribed as present day atmospheric pressure and present day salinity of the infiltrating water (for meteoric water S \approx 0). Note, that for F_{pr} = 0 (1) reduces to the UA-model.

As in the UA- and PR-models, Aeschbach-Hertig et al. (2000) assume that the gas-concentrations in the recharging water initially are in equilibrium with the atmosphere at atmospheric pressure and soil temperature and that bubbles of at-

	volume fraction	C_{A}^{*}/C_{B}^{*} at 5°C (-)	C_{A}^{*}/C_{B}^{*} at 10°C (-)	C_{A}^{*}/C_{B}^{*} at 20°C (-)	C_{A}^{*}/C_{B}^{*} at 30°C (-)	D_A/D_B at 5°C (-)	D_A/D_B at 10°C (-)	D_A/D_B at 20°C (-)	D_A/D_B at 30°C (-)
³ He/ ⁴ He	$1.384 \cdot 10^{-6}$	$1.3596 \cdot 10^{-6}$	$1.3602 \cdot 10^{-6}$	$1.3614 \cdot 10^{-6}$	$1.3625 \cdot 10^{-6}$	1.155	1.155	1.155	1.155
²⁰ Ne/ ²² Ne	9.8000	9.7804	9.7804	9.7804	9.7804	1.049	1.049	1.049	1.049
⁵⁶ Ar/ ⁴⁰ Ar Ne/Ar	$3.384 \cdot 10^{-3}$ $1.946 \cdot 10^{-3}$	$3.380 \cdot 10^{-3}$ $4.870 \cdot 10^{-4}$	$3.380 \cdot 10^{-3}$ $5.225 \cdot 10^{-4}$	$3.380 \cdot 10^{-3}$ $5.935 \cdot 10^{-4}$	$3.380 \cdot 10^{-3}$ $6.633 \cdot 10^{-4}$	1.054 1.596	1.054 1.573	1.054 1.532	1.054 1.494

Table 1b. Ratios of partial pressure in the atmosphere, of equilibirum concentration and of molecular diffusivity at different temperatures for selected noble gas isotopes and Ar and Ne.

mospheric air are trapped. However, instead of assuming complete dissolution of bubbles, they postulate that a reservoir of entrapped gas remains in the quasi-saturated zone. Water and entrapped gas are thought to form a closed system which equilibrates at hydrostatic pressure and surrounding soil temperature. This process changes the concentrations and the relative composition of noble gases in the groundwater and in the entrapped gas. The gas concentrations in the water resulting from the closed-system equilibration can be described by (CEmodel, Aeschbach-Hertig et al., 2000):

$$C_{i}(T,S,P,A,F) = C^{*}_{i}(T,S,P) + \frac{(1 - F_{cc})A_{cc}Z_{i}}{1 + F_{cc}A_{cc}Z_{i}/C^{*}_{i}(T,S,P)}$$
(2)

The parameter A_{ce} describes the initial STP-volume of dry air in the trapped gas per unit mass of water and the parameter F_{ce} describes the reduction of the volume of entrapped air A_{ce} due to partial dissolution and compression of the gas volume. The value of F_{ce} is restricted to the interval between 0 and 1, where $F_{ce} = 0$ implies unfractionated excess air and $F_{ce} = 1$ implies no gas excess. As in the PR-model, temperature T and the parameters determining the gas excess and its fractionation, A_{ce} and F_{ce} , are usually treated as fit parameters whereas P and S at recharge are prescribed.

The UA-model, assuming complete dissolution of the entrapped air without subsequent gas loss, corresponds to the limiting case of the PR- and CE-models with no fractionation $(F_{pr}=F_{ce}=0)$. In this case, A_{pr} in the PR-model and A_{ce} in the CE-model assume the same value and the predicted recharge temperature *T* is the same in both models. However, if the gas excess is fractionated compared to the atmospheric gas composition, predictions based on the PR- and the CE-model differ significantly.

According to the PR-model fractionation of the gas excess is solely due to differences in the molecular diffusivities between the gases:

$$\frac{\Delta C_A}{\Delta C_B} = \frac{z_A}{z_B} \cdot \exp\left(-F_{pr} \frac{D_A - D_B}{D_{Ne}}\right) \text{ for } A_{pr} \neq 0 \qquad (3)$$

where $\Delta C = (C-C^*)$ is the gas excess, and the ratio of the excess of gas A to the excess of gas B is a measure of fractionation. Excess gas only exists if $A_{pr} \neq 0$. Choosing A and B such that $D_A > D_B$, the ratio of the gas excess $\Delta C_A / \Delta C_B$ tends to 0 for $F_{pr} \rightarrow \infty$. If $F_{pr} \rightarrow 0$ excess air has atmospheric composition ($\Delta C_A / \Delta C_B = z_A / z_B$). The ratio of the dissolved gas concentrations C_A / C_B can range between 0 ($A_{pr} \rightarrow \infty$ and $F_{pr} \rightarrow \ln\{A_{pr}/C_A^*\} \cdot D_{Nc}/D_A$), the maximum of $z_A / z_B (A_{pr} \rightarrow \infty)$ and $F_{pr} = 0$), and C_A^* / C_B^* ($A_{pr} = 0$ or $F_{pr} \rightarrow \infty$; i.e., no gas excess

exists). Note that the limits given above are those for model Eqn. 1 but that the underlying physical concept breaks down at very large values of A_{pr} because the initial ratio of air to water volume becomes unreasonably large. Figure 1a demonstrates how the concentration ratio of ²⁰Ne to ²²Ne, C_{20Ne}/C_{22Ne} , depends on excess air and fractionation in the PR-model, i.e., on the parameters A_{pr} and F_{pr} . At given A_{pr} , C_{20Ne}/C_{22Ne} decreases with increasing F_{pr} until a minimum concentration ratio is reached and then increases again to finally approach atmospheric equilibrium at large F_{pr} . The larger the amount of initial excess air A_{pr} the lower the minimum of C_{20Ne}/C_{22Ne} .

In the CE-model, fractionation does not depend on molecular diffusivities but on the differences in the Henry coefficients between the gas species and the ratio of water volume to volume of entrapped air:

$$\frac{\Delta C_A}{\Delta C_B} = \frac{z_A}{z_B} \cdot \frac{C_A^*}{C_B^*} \cdot \frac{C_B^* + F_{ce}A_{ce}z_B}{C_A^* + F_{ce}A_{ce}z_A} \text{ for } F_{ce} \neq 1$$

and $A_{ce} \neq 0$ (4)

where the dependence on the Henry coefficients is implicitly contained in the equilibrium concentrations C_A^* and C_B^* (Aeschbach-Hertig et al., 1999) and the dependence on water volume and volume of entrapped air is contained in the parameters F_{ce} and A_{ce} . Eqn. 4 assumes $F_{ce} \neq 1$ and $A_{ce} \neq 0$ because only then excess gas exists. $\Delta C_A / \Delta C_B$ ranges between C_A^* / C_B^* $(A_{ce} \rightarrow \infty, F_{ce} \neq 0)$ and the ratio in unfractionated air z_A/z_B (F_{ce} = 0). In contrast to the PR-model, the ratio of the dissolved gas concentrations C_A/C_B in the CE-model cannot approach 0 but must range between C_A^*/C_B^* ($F_{ce} = 1$ or $A_{ce} = 0$) and z_A/z_B $(F_{ce} \rightarrow 0, A_{ce} \rightarrow \infty \text{ and } F_{ce}A_{ce} \rightarrow 0)$. Note that the physical concept underlying the CE-model breaks down for $A_{ce} \rightarrow \infty$. Fig. 1b shows how the concentration ratio C_{20Ne}/C_{22Ne} varies with A_{ce} and F_{ce} . At given A_{ce} , the concentration ratio C_{20Ne} C_{22Ne} decreases monotonically with the parameter F_{ce} and reaches C_A^*/C_B^* at $F_{ce} = 1$. The concentration ratio at $F_{ce} = 0$ increases with increasing A_{ce} but cannot exceed z_{20Ne}/z_{22Ne} .

Eqn. 3 and 4 imply that noble gases with Henry coefficients of similar magnitude but with very different molecular diffusivies should react much more sensitive to fractionation according to the PR-model than the CE-model and could be used to test which model is applicable. Because of the very large molecular diffusivity of He (Table 1), the model choice especially affects the predicted ratio between He and the heavier noble gases (Ne, Ar, Kr, Xe). This is particularly important in ${}^{3}\text{H}{}^{-3}\text{He}$ groundwater dating, where the ${}^{3}\text{He}$ contribution of atmospheric origin, ${}^{3}\text{He}_{atm}$ has to be known. In case of fractionated excess air ${}^{3}\text{He}_{atm}$ and consequently the ${}^{3}\text{H}{}^{-3}\text{He}$ water age derived depend strongly on the gas exchange model em-



Fig. 1. ²⁰Ne/²²Ne ratio as function of A_{pr} and F_{pr} for the PR-model (a) and A_{ce} and F_{ce} for the CE-model (b). Each line represents the variation of ²⁰Ne/²²Ne ratio with F_{pr} (a) and F_{ce} (b) at a fixed value of A, i.e., a fixed A_{pr} and A_{ce} , respectively. Conditions at recharge are assumed to be $T = 30^{\circ}$ C, p = 1 atm and S = 0. Note that the CE-model only allows F_{ce} to range between 0 and 1.

ployed. The presence of terrigenic and radiogenic/tritiogenic He sources prevents the use of He to distinguish between the PR- and the CE-model.

Isotopes of the the same noble gas have very similar Henry coefficients but can have significantly different molecular diffusivities (Table 1b). That is especially the case for the light noble gas isotopes ³He - ⁴He, ²⁰Ne - ²²Ne, and ³⁶Ar - ⁴⁰Ar. According to the above discussion, the isotopic ratios ³He/⁴He, ²⁰Ne/²²Ne and ³⁶Ar/⁴⁰Ar in the gas excess should remain between the atmospheric ratio and the ratio at atmospheric solubility equilibrium if fractionation occurs according to the CE-model, but might be significantly smaller if the fractionation depends on the differences in molecular diffusivities (PR-model). Because the ³He/⁴He ratio is affected by radiogenic/tritiogenic and terrigenic sources and the ³⁶Ar/⁴⁰Ar ratio is also influenced by terrigenic sources in aquifers with very large groundwater ages (e.g., Beyerle et al., 2000b; Torgersen et al., 1989), the ²⁰Ne/²²Ne ratio is the most reliable noble gas isotope ratio to test the fractionation models. The ³⁶Ar/³⁸Ar ratio could in principle also be used, although it is less sensitive to fractionation than the ²⁰Ne/²²Ne ratio, but to our knowledge it has never been measured in groundwater studies, due to the high experimental requirements (good mass resolution needed to separate ³⁸Ar from ⁴⁰Ar). Although molecular diffusivities differ much more between heavy noble gases and Ne than between ²⁰Ne and ²²Ne (Table 1), the ²⁰Ne/²²Ne ratio is best suited to distinguish between the PR- and the CE-model because its dependence on recharge temperature is negligible and the ²⁰Ne/²²Ne ratio can be measured with a much higher precision (typically 0.2%) than the concentrations of elemental noble gases (typically 1 to 2%).

2.2. Analysis of Noble Gas Data using the Software NOBLE

Noble gas data analysis with respect to recharge temperature, excess air and its fractionation is commonly performed by inverse fitting based on χ^2 -minimization, which takes the error of the measurements into account (Aeschbach-Hertig et al., 1999). The new fitting program Noble developed for this study extends the original fitting procedure of Aeschbach-Hertig et al. (1999) by incorporating the CE-model and including the possibility to consider not only elemental noble gas concentrations but also concentrations of noble gas isotopes and noble gas isotope ratios. Commonly the three unknown parameters $T, A_{\mu\nu}$ F_{pr} or T, A_{ce} , F_{ce} , respectively, are estimated from 4 observables, i.e., the concentration of Ne, Ar, Kr and Xe. Pressure P and salinity S at recharge are prescribed. Considering the ²⁰Ne/ ²²Ne and/or the ³⁶Ar/⁴⁰Ar ratios in addition to the elemental concentrations increases the number of observables and hence allows to estimate additional parameters (e.g., P if the location of recharge is unknown) or can be used to improve the constraint on the fit parameters.

Noble extends the original parameter set employed in Aeschbach-Hertig et al. (1999) by including optional parameters that describe the concentrations of terrigenic ⁴He and tritiogenic ³He, as well as the terrigenic ³He/⁴He ratio. An additional scaling parameter enables analysis based on the relative concentrations of the noble gases if the scaling of the absolute concentrations is unknown (e.g., unknown sample weight). A similar scaling technique has been used to interpret noble gas data from natural gas deposits where the degassing water volume is unknown (Ballentine et al., 1999). Furthermore, Noble supports ensemble fitting, i.e., the model parameters can be fitted not only to the gas concentrations of individual water samples separately but also to the concentrations from an ensemble of samples simultaneously. Thereby some of the fit parameters can be assumed to have the same value for all samples while other parameters may differ between samples. Propagation of the error of the measurements is performed by linear error propagation and optionally by a Monte-Carlo procedure to provide errors of the fit parameters. Both methods assume that measurement errors are normally distributed and independent. Noble calculates the probability $p(\chi^2)$ for the minimum χ^2 obtained from the fitting to be observed at the given number of freedoms. This provides the possibility to apply a χ^2 -test to the models. Noble provides the probability $p(\chi^2)$ for each water sample separately and for the entire data set $p(\chi^2_{set})$, where χ^2_{set} is the sum of the χ^2 values obtained for the samples of the data set. NOBLE will be made available on the Internet (http://www.eawag.ch/research_e/w+t/UI/ noblegasmethod.html).

3. STUDY AREA AND DATA

Details on the study area and an extended data set are given in Beyerle et al. (2002). In brief, groundwater samples were taken from the Continental Terminal (CT) aquifers located in south-western Niger, Africa, between 12.5 to 14.5° N and 2.5 to 4.5° E (Fig. 2). Within the CT formation three different aquifer compartments can be distinguished (CT3, CT2, CT1). CT3 is mostly unconfined, whereas CT2 and CT1 are confined. The Continental Terminal aquifer system is separated against the underlying Continental Intercalaire aquifer (CI) by massive low-permeable sediments mainly consisting of clays (Andrews et al., 1994). The altitude of the investigated area is almost constant increasing only slightly from 200 m to 300 m a.s.l. in north-easterly direction. The recharge area of the CT2 aquifer is located in the northern part of the study area and has an average altitude of 300 m a.s.l whereas the recharge areas of the CT1 and CI aquifers are further to the north at a mean altitude of \sim 350 to 400 m.

Groundwater samples were collected in April 1999 and June 2000 from drinking water wells either operated by manual pumps or by electric submersible pumps. In all samples, concentrations of the noble gas isotopes ³He, ⁴He, ²⁰Ne, ²²Ne, ³⁶Ar, ⁴⁰Ar, ⁸⁶Kr and ¹³⁶Xe were analysed according to the methods described in Beyerle et al. (2000a).

4. RESULTS AND DISCUSSION

4.1. Comparison of Gas Exchange Models

Figure 3 compares the model results from the PR- and the CE-model on recharge temperature for the CT aquifers derived by inverse fitting of the concentrations of Ne, Ar, Kr and Xe. Recharge temperatures are displayed as function of terrigenic ⁴He (⁴He_{ter}), which can be interpreted as a qualitative measure of groundwater residence time because ⁴He_{ter} (here of crustal origin) accumulates over time. ⁴He_{ter} is the difference between the measured ⁴He concentration and the concentration of ⁴He of atmospheric origin (⁴He_{atm}) determined from the PR- and the CE-model respectively. The concentration of atmospheric



Fig. 2. Map of the investigated area including sampling locations. The altitude of the area is almost constant showing a slight gradient in north-eastern direction from 200m to the CT2 recharge area at \sim 300m a.s.l. The CT1 and CI systems are recharged even further to the north-east or east at altitudes of \sim 350 to 400m. The depth of the unconfined groundwater table in the recharge zones varies between 5 and 20m in the Dallols (dry, sand filled riverbeds) and between 20 and 60m elsewhere.

origin is defined as the sum of two contributions: the atmospheric equilibrium concentration at T, S and P and the concentration due to fractionated excess air.

The model assuming unfractionated excess air has to be rejected according to the χ^2 -test, making it necessary to account for fractionation. Both the PR-model and the CE-model predict Ne, Ar, Kr and Xe concentrations compatible with the data. The PR-model (open symbols, Fig. 3) yields systematically higher recharge temperatures than the CE-model (solid symbols, Fig. 3), on average by 1.5° C and for the 3 oldest samples even by $\sim 4^{\circ}$ C. The difference in recharge temperature between young samples (${}^{4}\text{He}_{\text{ter}} < 1.5 \cdot 10^{-7} \text{ cm}^{3}\text{STPg}^{-1}$; n=17) and intermediate samples ($1.5 \cdot 10^{-7} \text{ cm}^{3}\text{STPg}^{-1} < {}^{4}\text{He}_{\text{ter}} < 1.5 \cdot 10^{-6} \text{ cm}^{3}\text{STPg}^{-1}$; n=7) is about the same for the PR-model ($\Delta T = 4.0^{\circ}$ C ± 1.8° C) and the CE-model (4.8° C ± 1.3° C). A thorough discussion of paleo-climatic conditions in Niger based on

a more comprehensive data set on tracers from the CT Aquifers is given elsewhere (Beyerle et al., 2002). Here we focus on the use of Ne isotopes to distinguish between the two models.

Figure 4 compares the measured ${}^{20}\text{Ne}/{}^{22}\text{Ne}$ (Fig. 4a) and ${}^{36}\text{Ar}/{}^{40}\text{Ar}$ ratios (Fig. 4b) with those predicted from the PRmodel (open symbols) and the CE-model (solid symbols). The model parameters were derived by inverse fitting of Ne, Ar, Kr and Xe concentrations. The agreement between data and model prediction is significantly better for the CE-model than for the PR-model. Within error almost all measured ${}^{20}\text{Ne}/{}^{22}\text{Ne}$ ratios ((${}^{20}\text{Ne}/{}^{22}\text{Ne})_{m}$) fall in the range between the isotopic ratio in the atmosphere (Ozima and Podosek, 1983) (Fig. 4a dashed lines, $z_{20\text{Ne}}/z_{22\text{Ne}}$) and the isotopic ratio at atmospheric equilibrium (Beyerle et al., 2000a) (Fig. 4a dotted lines, ${}^{20}\text{Ne}^{*}/$ ${}^{22}\text{Ne}^{*}$). Considering the standard deviation of all data, the mean of the measured ${}^{20}\text{Ne}/{}^{22}\text{Ne}$ and ${}^{36}\text{Ar}/{}^{40}\text{Ar}$ ratios are consistent



Fig 3. Recharge temperature as function of ⁴He_{ter} determined by inverse fitting based on the noble gas concentrations Ne, Ar, Kr and Xe using the CE-model (solid symbols) and the PR-model (open symbols). Fit parameters are T, A_{ce} and F_{ce} in case of the CE-model and T, A_{pr} and F_{pr} in case of the PR-model. ⁴He_{ter} can be interpreted as a qualitative groundwater age, because in groundwaters terrigenic helium accumulates over time.

with a value between the corresponding ratios in the atmosphere and at atmospheric equilibrium (Table 2).

As discussed above, fractionation described by the CEmodel always predicts isotopic ratios in the range between the values for air and atmospheric equilibrium. The PR-model however may predict isotopic ratios below this range as is demonstrated in the case of the CT Aquifers (Fig. 4 open symbols, Table 2).

The constraint on the fit-parameters *T*, *A* and *F* can be increased if the measured ²⁰Ne/²²Ne ratio is included as an observable in addition to the noble gas concentrations of Ne, Ar, Kr, and Xe. Using this extended data set, the CE-model yields a good fit ($p(\chi^2_{set}=67) \approx 11\%$), whereas the PR-model has to be rejected ($p(\chi^2_{set}=440) << 1\%$). This result reflects the fact that the measured ²⁰Ne/²²Ne ratios agree better with the predictions according to the CE-model than with the predictions according to the PR-model.

The best fit values for the parameters of the CE-model derived for the extended data set with the ²⁰Ne/²²Ne ratio included are essentially the same as those derived from the elemental concentrations of Ne, Ar, Kr and Xe only. In contrast, including the ²⁰Ne/²²Ne ratio reduces the best fit values for F_{pr} of the PR-model by a factor of 10, resulting in F_{pr} ranging from 0 to 0.27 with a mean of $F_{pr} = 0.1 \pm 0.1$. This implies that the predicted fractionation is small and that the PR-model essentially reduces to the UA-model. The constraint provided by the measured Ne isotopes ratios rules out significant diffusive gas loss.

Because the PR-model is incompatible with the data when considering measured ²⁰Ne/²²Ne ratios, only the CE-model adequately describes the gas exchange in the CT aquifers. Thus in case of the CT aquifers the parameters describing recharge temperature, gas excess and its fractionation derived by inverse fitting of Ne, Ar, Kr, Xe and the ²⁰Ne/²²Ne ratio using the CE-model should be employed in the discussion of paleoclimatic change and for the calculation of ³He_{atm} and ⁴He_{atm} for dating purposes.

Observed ²⁰Ne/²²Ne ratios fall within the range between the ratio in the atmosphere and the ratio at atmospheric equilibrium not only in the CT aquifers of Niger but also in other aquifers where fractionated excess gas has been reported. Aeschbach-Hertig et al. (2000) analysed four data sets, but Ne isotope data are unfortunately available only for two of them (Belgium and Oman). In both cases no significant fractionation of the Ne isotopes was observed, e.g., in northern Oman (Weyhenmeyer et al., 2000) the ²⁰Ne/²²Ne ratio measured in 9 samples ranges between 9.763 and 9.865 with a mean of 9.795 \pm 0.029. This supports the applicability of the CE-model in noble gas based paleo-studies and for groundwater age dating. However, it does not necessarily exclude the PR-model. The PR- and the CEmodel only describe the mean effect of gas exchange on noble gases and do not resolve details on the interaction between gas excess, entrapped air and recharging groundwater. Therefore, it is still unclear which environmental conditions lead to the noble gas pattern described by the PR-model and which result in the noble gas pattern described by the CE-model. Thus,



Fig. 4. Comparison of measured and predicted ²⁰Ne/²²Ne and ³⁶Ar/⁴⁰Ar ratios. Predictions were calculated from the model parameters describing recharge temperature, excess air and its fractionation derived with the CE-model (solid symbols) and PR-model (open symbols), respectively. Model parameters were obtained by inverse fitting to the concentrations of Ne, Ar, Kr and Xe.

Table 2. Mean measured and predicted ²⁰Ne/²²Ne and ³⁶Ar/⁴⁰Ar ratios in comparison with the corresponding ratios in the atmosphere and in atmospheric equilibrium.

	²⁰ Ne/ ²² Ne	³⁶ Ar/ ⁴⁰ Ar (10 ⁻³)
Mean of measurements	9.798 ± 0.024	3.390 ± 0.008
Mean of predicted ratios using the PR-model	9.604 ± 0.095	3.350 ± 0.020
Mean of predicted ratios using the CE-model	9.785 ± 0.002	3.381 ± 0.0003
Isotope ratios in the atmosphere (Ozima and Podosek, 1983)	9.8000	3.384
Isotope ratios in atmospheric equilibrium (Beyerle et al.,	9.7804	3.380
2000a)		

whether the PR- or the CE-model should be applied in a specific aquifer can be judged only from data on concentrations of noble gases and their isotopes.

4.2. Dating Applications

Groundwater dating with the ³H-³He method requires estimation of tritiogenic ³He (³He_{tri}) by subtracting ³He_{atm} and terrigenic ³He (³He_{ter}) from the measured ³He (³He_m) concentrations. Assuming that ³He_{atm}, ⁴He_{atm} and the ³He/⁴He ratio of the terrigenic component (³He/⁴He)_{ter} is known, ³He_{tri} can be calculated from the balance of ⁴He and ³He (e.g., Schlosser et al., 1989):

$$He_{ter} = {}^{4}He_{m} - {}^{4}He_{atm} {}^{3}He_{tri} = {}^{3}He_{m} - {}^{3}He_{atm} - {}^{4}He_{ter} \cdot \left(\frac{{}^{3}He}{{}^{4}He}\right)_{ter}$$
(5)

4.2.1. Estimation of ${}^{3}He_{atm}$ and ${}^{4}He_{atm}$

The ³He and ⁴He components of atmospheric origin can be estimated from the ³He and ⁴He concentrations in the atmosphere (Ozima and Podosek, 1983) together with the ³He and ⁴He solubilities (Benson and Krause, 1980; Weiss, 1971) by employing an appropriate gas exchange model (e.g., the PR- or the CE-model) where model parameters describing gas excess and fractionation have been estimated by inverse fitting of noble gas concentrations. In case of the CT aquifers in Niger estimated ³He_{atm} and ⁴He_{atm} differ substantially between the PR- and CE- model if the inverse fitting procedure is based only on the elemental concentrations of Ne, Ar, Kr and Xe (Fig. 5a, Table 3).

If the measured ²⁰Ne/²²Ne ratio is included as additional observable, ³He_{atm} and ⁴He_{atm} predicted with the PR-model increase substantially and closely agree with predictions assuming unfractionated excess air (Table 3). Predictions of ³He_{atm} and ⁴He_{atm} based on the CE-model are independent of whether the (²⁰Ne/²²Ne)_m is included in the parameter estimation or not (see Table 3). Because only the CE-model is compatible with the measured elemental noble gas concentrations and the ²⁰Ne/²²Ne ratio (see above) the ³He_{atm} and ⁴He_{atm} predicted with the CE-model are assumed to correspond to the true values.



Fig. 5. ${}^{3}\text{He}_{atm}$ and ${}^{3}\text{H}$ - ${}^{3}\text{He}$ water ages for young groundwaters of the CT aquifers. Concentrations of ${}^{3}\text{He}_{atm}$ are shown for all samples with $\text{He}_{ter} < 1.5 \cdot 10^{-7} \text{ cm}^{3}\text{STPg}^{-1}$, which all infiltrated at similar temperatures of $\sim 32^{\circ}\text{C}$ (see Fig. 2a). ${}^{3}\text{H}$ - ${}^{3}\text{He}$ water ages are shown for all samples in which ${}^{3}\text{H}$ has been detected and ${}^{3}\text{He}_{tri}$ is significantly larger than zero (b). Results from the PR-model and the CE-model based on Ne, Ar, Kr and Xe are presented and compared to results from simplified procedures which are based on Ne only and on the combination of Ne, Ar and ${}^{20}\text{Ne}/{}^{22}\text{Ne}$. The ${}^{3}\text{H}$ - ${}^{3}\text{He}$ water ages were calculated assuming $({}^{3}\text{He}/{}^{4}\text{He})_{ter} = (2.2 \pm 0.1) \cdot 10^{-8}$ except in the case of the PR-model where $({}^{3}\text{He}/{}^{4}\text{He})_{ter} = (3.0 \pm 0.1) \cdot 10^{-8}$ was used.

4.2.2. Estimation of $({}^{3}He/{}^{4}He)_{ter}$ and natural (pre-bomb) Tritium

By restriction to samples that are not influenced by bomb tritium, the values of $({}^{3}\text{He}/{}^{4}\text{He})_{ter}$ and the ${}^{3}\text{He}_{tri}$ of prebomb origin can be estimated by inverse fitting utilising the ensemble fitting option of Noble. To this end, noble gas concentrations and isotope ratios from all intermediate and old groundwater samples (${}^{4}\text{He}_{ter} > 1.5 \cdot 10^{-7} \text{ cm}^{3}\text{STPg}^{-1}$; 10 samples) were compiled into a data ensemble. The fit-parameters *T*, *A* and *F* determining the noble gas concentration of atmospheric origin and the fit parameter describing the concentration of ${}^{4}\text{He}_{ter}$ are assumed to be different for each water sample. (${}^{3}\text{He}/{}^{4}\text{He})_{ter}$ and prebomb ${}^{3}\text{He}_{tri}$ are assumed to be the same for all water samples, i.e., for the data ensemble.

Table 3. The effect of model choice and availability of data on χ^2_{set} and predicted ³He_{atm}. Data originate from young goundwater samples (n=17) of the CT aquifers in Niger. ³He_{atm} predicted from the CE-model based on Ne, Ar, Kr, Xe and ²⁰Ne/²²Ne serves as reference. The degrees of freedom

	Decomptors	Parameters prescribed			Deviation of ³ He _{atm} from the best estimate (%)	
Data considered	fitted		$\chi^2_{ m set}$	$\nu_{\rm set}$	minimum	maximum
CE-model						
Ne, Ar, Kr, Xe, ²⁰ Ne/ ²² Ne	T, A_{ad}, F_{ad}		38	34	0	0
Ne, Ar, Kr, Xe	T, A _{ac} , F _{ac}		8	17	-0.06	+0.04
Ne, Ar	A _{ce} , F _{ce}	$T = 32^{\circ}\mathrm{C}$	0	0	-0.2	+0.2
Ne, 20 Ne/ 22 Ne	A_{ad}^{ce} F_{ad}^{ce}	$T = 32^{\circ}\mathrm{C}$	24	0	-3.9	+3.2
Ne, Ar, 20 Ne/ 22 Ne	A_{ce}^{ce} , F_{ce}^{ce}	$T = 32^{\circ}\mathrm{C}$	29	17	-0.2	+0.2
Ne, Ar, 20 Ne/ 22 Ne	A_{ad}^{ce} F_{ad}^{ce}	$T = 30^{\circ}\mathrm{C}$	50	17	-0.8	+0.2
Ne, Ar, 20 Ne/ 22 Ne	A_{ad}^{ce} F_{ad}^{ce}	$T = 35^{\circ}\mathrm{C}$	47	17	-0.09	+0.8
PR-model	ce ce					
Ne, Ar, Kr, Xe, ²⁰ Ne/ ²² Ne	T, A_{ca}, F_{ca}		176	34	-5.4	+1.7
Ne, Ar, Kr, Xe	T, A_{ce}, F_{ce}		5	17	-27.8	-6.2
Ne, Ar	A _{art} F _{ar}	$T = 32^{\circ}\mathrm{C}$	0	0	-36.3	-8.3
Ne, 20 Ne/ 22 Ne	A_{ad}^{ce} F_{ad}^{ce}	$T = 32^{\circ}\mathrm{C}$	21	0	-2.8	+3.2
Ne, Ar, 20 Ne/ 22 Ne	A_{ad}^{ce} F_{ad}^{ce}	$T = 32^{\circ}\mathrm{C}$	570	17	-14.8	+2.3
UA-model	ce ce					
Ne, Ar, Kr, Xe, ²⁰ Ne/ ²² Ne	T, A	F = 0	191	51	+0.4	+2.6
Ne, Ar, Kr, Xe	T, A	F = 0	164	34	+0.4	+2.6
Ne	À	$T = 32^{\circ}C; F = 0$	0	0	+0.7	+3.2
Ne, Ar	Α	$T = 32^{\circ}C; F = 0$	859	17	+1.0	+7.9
Ne, 20 Ne/ 22 Ne	Α	$T = 32^{\circ}C; F = 0$	27	17	+0.7	+3.2
Ne, Ar, ²⁰ Ne/ ²² Ne	Α	$T = 32^{\circ}\text{C}; F = 0$	886	34	+1.0	+8.5

First we consider the data set consisting of the measured elemental concentrations of He, Ne, Ar, Kr, and Xe and the ³He/⁴He ratio for each of the 10 samples. Altogether 60 independent measurements are available for the inverse fitting of 42 fit parameters leaving 18 degrees of freedom. For this data set the CE-model gives $({}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}} = (2.2 \pm 0.1) \cdot 10^{-8}$ and ${}^{3}\text{He}_{\text{tri}} = 0.2 \pm 0.3$ TU (1 cm³STPg⁻¹ $\triangleq 4.019 \cdot 10^{14}$ TU). The value of ³He_{tri} corresponds to the concentration of prebomb tritium at the groundwater table. Prebomb ³H close to zero is reasonable because radioactive decay during the water transport through the large unsaturated zone of the CT aquifers substantially reduces the tritium concentration reaching the saturated zone (Brennwald et al., 2001). ⁴He_{ter} is different for each sample and can be used as a qualitative groundwater age as in Figure 3. If the PR-model is applied instead of the CE-model one obtains $({}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}} = (3.0 \pm 0.1) \cdot 10^{-8} \text{ and } {}^{3}\text{He}_{\text{tri}} = 11 \pm 1 \text{ TU. Both}$ values are significantly larger than those predicted by the CE-model. Including the ²⁰Ne/²²Ne ratios in the data set, 70 observables are available for the 42 fit parameters leaving 28 degrees of freedom. For the CE-model the predicted values of $({}^{3}\text{He}/{}^{4}\text{He})_{ter}$ and ${}^{3}\text{He}_{tri}$ are the same as above. In case of the PR-model $({}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}} = (2.1 \pm 0.2) \cdot 10^{-8}$ and ${}^{3}\text{He}_{\text{tri}} = 3 \pm 1$ TU. Both values are significantly smaller than those predicted without the ²⁰Ne/²²Ne ratios and closer to the results of the CE-model.

The parameters describing $({}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}}$, ${}^{4}\text{He}_{\text{ter}}$ and ${}^{3}\text{He}_{\text{tri}}$ are only affected by concentrations of ${}^{3}\text{He}$ and ${}^{4}\text{He}$, whereas the parameters *T*, *A* and *F* are affected by the concentrations of all noble gas isotopes. Hence, one can prescribe the parameters *T*, A_{ce} , and F_{ce} using the results from inverse fitting of Ne, Ar, Kr and Xe, and fit only the parameters describing $({}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}}$, ${}^{4}\text{He}_{\text{ter}}$ and ${}^{3}\text{He}_{\text{tri}}$ to the measured He concentration and ${}^{3}\text{He}/{}^{4}\text{He}$ ratio. This procedure involves 20 measurements and 12 fit parameters, i.e., 8 degrees of freedom. It underestimates the errors of the values for $({}^{3}\text{He}/{}^{4}\text{He})_{\text{terr}}$, ${}^{4}\text{He}_{\text{ter}}$ and ${}^{3}\text{He}_{\text{tri}}$ because the uncertainties of *T*, A_{ce} , F_{ce} are not propagated. The results for the two procedures are significantly different only in the case of the PR-model. However, for neither of the two procedures and models the model results are consistent with the data according to a χ^{2} test, indicating that the model assumption that (${}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}}$ and ${}^{3}\text{He}_{\text{tri}}$ are the same for all samples is too simplified to adequately describe the data set. Nevertheless the calculated values of (${}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}}$ and ${}^{3}\text{He}_{\text{tri}}$ are used as best estimates.

Figure 6a displays the predicted concentration of non-atmospheric ³He, i.e., the sum of ³He_{ter} and ³He_{tri}, versus the predicted ⁴He_{ter} for the PR- and the CE-model obtained from inverse fitting of *T*, A_{pr} , and F_{pr} or *T*, A_{ce} and F_{ce} , respectively, to the elemental concentrations of Ne, Ar, Kr and Xe. The regression lines give in case of the CE-model (³He/⁴He)_{ter} = (2.2 ± 0.1) · 10⁻⁸ and ³He_{tri} = 0.4 ± 0.2 TU and in case of the PR-model (³He/⁴He)_{ter} = (3.4 ± 0.1) · 10⁻⁸ and ³He_{tri} = 12 ± 1 TU. Fitting regression lines corresponds to the second approach taken above and confirms the results provided by the ensemble fitting using Noble.

In previous studies (e.g., Aeschbach-Hertig et al., 2000; Kipfer et al., 1994; Kipfer et al., 1996) (${}^{3}\text{He}/{}^{4}\text{He})_{ter}$ has been derived from a regression line in a three isotope plot ${}^{3}\text{He}/{}^{4}$ He vs Ne/ 4 He (Fig. 6b). Such a procedure assumes that all measured ratios can be interpreted as a mixture between a terrigenic component with constant (${}^{3}\text{He}/{}^{4}\text{He})_{ter}$ and (Ne/ He)_{ter} = 0, and a combined atmospheric/tritiogenic component with constant composition given by (${}^{3}\text{He}_{atm} + {}^{3}\text{He}_{tri}$)/ ${}^{4}\text{He}_{atm}$ and Ne_{atm}/He_{atm}. Although these assumptions are not strictly fulfilled, the variations in the atmospheric/tritiogenic endmember (due to varying *T*, *A*, *F*, and ${}^{3}\text{He}_{tri}$) are relatively small compared to the effect of the terrigenic component. Therefore this approach yields a useful estimate of (${}^{3}\text{He}/$

in the fits are given for the entire data set by v_{set} .



Fig. 6. Different methods to estimate $({}^{3}\text{He}/{}^{4}\text{He})_{ter}$.(a) Non-atmospheric ${}^{3}\text{He}$ versus non-atmospheric ${}^{4}\text{He}$ for intermediate and old groundwater samples (He_{ter} > 1.5 \cdot 10⁻⁷ cm³STPg⁻¹) from the CT aquifers. Assuming a constant prebomb tritium and a constant (${}^{3}\text{He}/{}^{4}\text{He})_{ter}$ the intercept of the regression line corresponds to ${}^{3}\text{He}_{tri}$ and the slope corresponds to (${}^{3}\text{He}/{}^{4}\text{He})_{ter}$. Based on the concentrations of Ne, Ar, Kr, and Xe, ${}^{3}\text{He}_{m}$ - ${}^{3}\text{He}_{atm}$ and ${}^{4}\text{He}_{m}$ - ${}^{4}\text{He}_{atm}$ are determined using the PR-model (open circles) and the CE-model (solid circles), respectively.(b) Measured ${}^{3}\text{He}/{}^{4}\text{He}$ ratio versus the measured concentration ratio of Ne to He. If the He concentration approaches infinity, the Ne/He ratio approaches zero and the atmospheric gas contribution becomes negligible compared to terrigenic contributions. Thus the intercept of the regression line in Fig. 5b can be interpreted to correspond to (${}^{3}\text{He}/{}^{4}\text{He})_{ter}$.

 ${}^{4}\text{He})_{\text{ter}} = (2.3 \pm 0.1) \cdot 10^{-8}$. This estimate agrees with the value found by the CE-model but is significantly lower than the value estimated from the PR-model based on the data set without the ${}^{20}\text{Ne}/{}^{22}\text{Ne}$ ratio.

4.2.3. Estimation of water age of young groundwater samples

The estimated values of $({}^{3}\text{He}/{}^{4}\text{He})_{\text{ter}}$, ${}^{3}\text{He}_{atm}$ and ${}^{4}\text{He}_{atm}$ from the different models can be employed to estimate ${}^{3}\text{He}_{rri}$ in young groundwater samples containing tritium. The resulting ${}^{3}\text{He}_{\text{tri}}$ differs between models and so does the ${}^{3}\text{H}$ - ${}^{3}\text{He}$ water age τ (Tolstikhin and Kamenskiy, 1969; Torgersen et al., 1979):

$$\tau = \frac{1}{\lambda} \ln \left(1 + \frac{{}^{3}He_{tri}}{{}^{3}H} \right)$$
(6)

where $\lambda = 0.05626 \text{ yr}^{-1}$ (Lucas and Unterweger, 2000). Depending on the choice of the gas exchange model, ³H-³He water ages differ by up to a factor of 2 (Fig. 5b). Shown are the water ages for all samples in which ³H has been detected and ³He_{tri} is significantly larger than zero. The water ages predicted by the PR-model appear unrealistically large. They suggest that around 1930 the tritium concentration at recharge $({}^{3}H_{rech} =$ ${}^{3}\text{H}_{m}$ + ${}^{3}\text{He}_{tri}$), i.e., at the bottom of the 30 m thick unsaturated zone, were up to 10 TU, which seems unrealistic. A consistency test of the ³H-³He water ages based on the comparison of ³H_{rech} with the historic ³H concentration at recharge (Aeschbach-Hertig et al., 1998; Stute et al., 1997) is not practicable in the CT-aquifers because the ³H at recharge is strongly affected by the transport through the unsaturated zone. Hence, only the inclusion of the Ne isotopes in the analysis provides an objective, i.e., data-based, criterion to identify the ³H-³He water ages predicted by the CE-model to be more reliable than those predicted by the PR-model.

4.2.4. Simplified procedures to estimate He_{atm}

In many studies the dating procedure outlined above is not applicable because measurements of heavy noble gases are not available. Ne can be used to estimate excess air even if the recharge temperature is only approximately known, because the equilibrium concentration Ne^{*} only weakly depends on temperature (Table 1). Usually (e.g., Aeschbach-Hertig et al., 1998; Ekwurzel et al., 1994; Schlosser et al., 1989; Torgersen et al., 1979) the Ne excess ($\Delta Ne = Ne_m$ -Ne^{*}) has been used to calculate the atmospheric He excess from $\Delta He = \Delta Ne \cdot (z_{He}/z_{Ne})$, i.e., assuming unfractionated excess air (UA-model). However, if the excess air is fractionated, its He/Ne ratio can be significantly lower than z_{He}/z_{Ne} (Holocher et al., 2001), particularly in the case of the PR-model.

The equivalent application of the UA-model to the Ne data from young groundwater samples (${}^{4}\text{He}_{ter}$ < 1.5 \cdot 10⁻⁷ cm³STPg⁻¹) of the CT aquifers, assuming the recharge temperature T to correspond to the annual mean soil temperature of ~32°C (CESBIO/ORSTOM/CNES.HAPEX SAHEL Information System, 1996), results in predicted concentrations of ³He_{atm} that are 0.7 to 3.2% larger than those predicted by the CE-model using the concentrations of all noble gases and the 20 Ne/ 22 Ne ratio (Fig. 5a, Table 3). This demonstrates that the estimation of ${}^{3}\text{He}_{atm}$ based on ΔNe and the assumption of unfractionated excess air provides a reasonable approximation if the gas excess is not strongly fractionated and fractionation is not of PR-model type. However, the information contained in the Ne concentration is not sufficient to test the validity of the assumption that fractionation is small or to distinguish between the different models.

It is possible to estimate the effect of fractionation on ⁴He_{atm} and ³He_{atm} if the ²⁰Ne/²²Ne ratio is considered in addition to the elemental Ne concentration. Using these two constraints and assuming that recharge temperature is approximately known, the model Eqn. 1 and 2 can be solved for the parameters A_{pr} and F_{pr} or A_{ce} and F_{ce} , respectively. With the derived model parameters, ⁴He_{atm} and ³He_{atm} can be calculated from Eqn. 1 and 2, respectively. Equivalently, Noble can be employed to fit the parameters A_{pr} and F_{pr} or A_{ce} and F_{ce} , respectively, based on the measured Ne concentration, the ²⁰Ne/²²Ne ratios and a prescribed recharge temperature.

Application of this method to the Ne concentrations and ²⁰Ne/²²Ne ratios from the CT aquifers provides ³He_{atm} and ⁴He_{atm} which are very similar for all three models (PR-, CEand UA-model). The reason is that the inverse fitting based only on Ne concentrations and the ²⁰Ne/²²Ne ratio suggests that fractionation is small. The deviation between the ³He_{atm} predicted using the reduced data set and the best estimate of ³He_{atm} predicted using all available data ranges between -3.9% and 3.2% for all young groundwater samples with ${}^{4}\text{He}_{\text{ter}} < 1.5 \cdot 10^{-7} \text{ cm}^{3}\text{STPg}^{-1}$ (see Table 3). Although a clear decision between UA-, CE- and PR-model is impossible on the basis of Ne concentration and ²⁰Ne/²²Ne ratio, the range of possible ³He_{atm} can be estimated and strong fractionation of PR-model type can be distinguished from fractionation according to the CE-model or from unfractionated excess air as in the UA-model (Table 3).

Compared to the PR-model, the He-Ne fractionation according to the CE-model is not very different from unfractionated excess air (UA-model). Hence for dating studies the distinction between the CE- and the UA-model is not as critical as the distinction between the CE-and the PR-model. Nevertheless, in the case of samples with very young age, for which the difference between total ³He and ³He_{atm} is small, and in the case of a large contribution of excess gas, the prediction of ³He_{tri} and water age may differ significantly depending on whether CE- or UA-model is applied. Since the UA-model corresponds to the CE-model with $F_{ce} = 0$, the distinction between CE- and UA-model essentially is the problem of determining the parameters A_{ce} and F_{ce} of the CE-model from a limited data set on noble gases and noble gas isotopes. The $^{20}\mbox{Ne}/^{22}\mbox{Ne}$ ratio is not suitable to determine whether $F_{ce} = 0$ or not, because the solubilities of the Ne isotopes are very similar such that the ²⁰Ne/²²Ne ratio in atmospheric equilibrium is approximately the same as in the atmosphere. However, estimation of A_{ce} and F_{ce} could be based on the Ar concentration together with the Ne concentration and the 20Ne/22Ne ratio. The Ne/Ar ratio in atmospheric equilibrium differs significantly from that in the atmosphere (Table 2) and Ar concentrations can be measured comparatively easy and are less dependent on recharge temperature than the concentrations of the heavy noble gases Kr and Xe.

In case of the Niger data, the constraint by Ne and Ar concentrations together with the ²⁰Ne/²²Ne ratios is sufficient to rule out the UA- and PR-models, whereas the predictions according to the CE-model are compatible with the data from the young groundwater samples according to a χ^2 -test (p($\chi^2_{set}=29$) > 3%). Considering the groundwater samples individually, predictions according to the CE-model are consistent with the data of all samples (p(χ^2)>1% for all 27

samples) while the predictions of the PR- and the UA-model are acceptable only for 2 and 3 samples, respectively. Note that including the ²⁰Ne/²²Ne ratio is required to distinguish between the PR- and CE-model. Fitting based on Ne and Ar concentrations alone only allows rejection of the UA-model (Table 3).

The parameters A_{ce} and F_{ce} estimated from inverse fitting based on Ne and Ar concentrations and the ²⁰Ne/²²Ne ratio can be employed to determine the concentration of ³He_{atm} and ⁴He_{atm}. The ³He_{atm} concentrations agree within \pm 0.2% with the reference value obtained from the concentrations of Ne, Ar, Kr, Xe and the ²⁰Ne/²²Ne ratio (Table 3). In Figure 5a the predicted ³He_{atm} (crosses) is compared to the predictions using the UA-model based only on Ne (triangles) and to the predictions using the CE- and the PR- models based on concentrations of Ne, Ar, Kr and Xe (solid and open circles, respectively).

The prediction of ³He_{atm} based on the concentrations of Ne, Ar and the ²⁰Ne/²²Ne ratio only weakly depends on *T*. For recharge temperatures between 30°C and 35°C ³He_{atm} deviates from the best value by less than 1% for the young groundwater samples (Table 3). The χ^2_{set} for the fits with T = 30°C and T = 35°C are larger than for T = 32°C, indicating that the latter is the most appropriate value for the recharge temperature, in agreement with the the soil temperature and the noble gas temperatures derived from all noble gas concentrations using the CE-model.

³H-³He water ages calculated from the ³He_{atm} and ⁴He_{atm} obtained from the simplified procedures discussed above agree well with the ³H-³He water ages derived from the entire data set using the CE-model (Fig. 5b). Water ages derived using the usual He correction based only on Δ Ne are typically 1 to 2 yr younger than the best estimates. This is particularly important in studies concerned with dating in very young groundwaters. In the samples 19 and 30 ³H and ³He_{tri} are very small, such that small uncertainties in ³He_{atm} lead to large errors in the ³H-³He water age. Water ages based on Ne, Ar and ²⁰Ne/²²Ne agree with the best estimates within error and in most cases the ³H-³He water ages deviate by less than 0.2 yr.

5. CONCLUSIONS

Concentrations of noble gas isotopes carry information on gas partitioning during groundwater recharge which might not be available from the elemental noble gas concentrations of Ne, Ar, Kr, and Xe. Specifically, in the CT-aquifers of Niger a decision between the PR- and the CE-model is impossible based only on the elemental concentrations. If however measurements of the ²⁰Ne/²²Ne ratio are included in the analysis, the PR-model has to be rejected according to a χ^2 -test and only the CE-model is compatible with the entire data set. This finding is also confirmed by the ³⁶Ar/⁴⁰Ar ratio, which is incompatible with the PR-model. Thus the concentrations of noble gas isotopes allow a decision between PR- and CE-model which influences the prediction of recharge temperature, initial amount of excess air, and fractionation and hence affects the interpretation of the data with respect to paleoclimatic conditions.

The distinction between PR-, CE- and UA-model is also important in ³H-³He dating. In groundwaters with fractionated gas excess, ³He_{atm} required to calculate ³He_{tri} strongly depends on the fractionation model employed. Commonly the excess gas contributing to ${}^{3}\text{He}_{atm}$ is estimated from ΔNe assuming unfractionated excess air (UA-model). The effect of fractionation can only be assessed by including additional experimental information. Combination of the Ne concentration with the ²⁰Ne/²²Ne ratio allows estimation of the range of possible values of ³He_{atm}, thus providing the range of possible water ages. However, this procedure is not suitable to distinguish between PR-, CE- and UA-model if fractionation is small, i.e., if the ²⁰Ne/²²Ne is between the value in the atmosphere and the value at atmospheric equilibrium. Extending the data set consisting of Ne concentration and ²⁰Ne/²²Ne ratio by the concentration of Ar enables a rigorous distinction between models and a reliable estimation of ³He_{atm} and ⁴He_{atm} and of ³H-³He water ages.

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