Hydrol. Earth Syst. Sci. Discuss., 8, 4559–4581, 2011 www.hydrol-earth-syst-sci-discuss.net/8/4559/2011/ doi:10.5194/hessd-8-4559-2011 © Author(s) 2011. CC Attribution 3.0 License.



This discussion paper is/has been under review for the journal Hydrology and Earth System Sciences (HESS). Please refer to the corresponding final paper in HESS if available.

Classification of thermal waters based on their inorganic fingerprint and hydrogeothermal modelling

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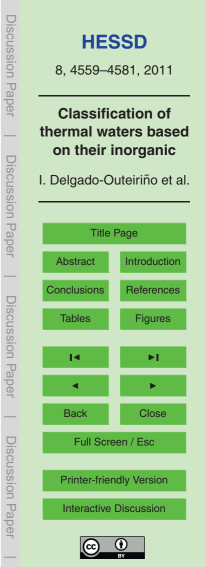
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Received: 29 March 2011 - Accepted: 15 April 2011 - Published: 5 May 2011

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Published by Copernicus Publications on behalf of the European Geosciences Union.



Abstract

Hydrothermal features in Galicia have been used since ancient times for therapeutic purposes. A characterization of these thermal waters was carried out in order to understand their behaviour based on inorganic pattern and water-rock interaction
mechanisms. In this way 15 thermal water samples were collected in the same hydrographical system. The results of the hydrogeochemistry analysis showed one main water family of bicarbonate type sodium waters, typical in the post-orogenic basins of Galicia. Principal component analysis (PCA) and partial lest squared (PLS) clustered the selected thermal waters in two groups, regarding to their chemical composition.
This classification agreed with the results obtained by the use of geothermometers and the hydrogeochemical modelling. The first included thermal samples that could be in contact with surface waters and therefore, their residence time in the reservoir and

in contact with surface waters and therefore, their residence time in the reservoir and their water-rock interaction would be less important than for the thermal waters of the second group.

15 **1** Introduction

20

Galicia, in northwest Spain and with an area of 29574 km², is bordered by Portugal to the south, the Spanish regions of Castile and León and Asturias to the east, the Atlantic Ocean to the west, and the Bay of Biscay to the north. Galicia was affected by the hercynianorogeny and in this region, materials of Proterozoic and the Palaeozoic outcrops are affected by major faults is Hesperian massif has emerged since the end of the Paleozoic and erosion has the panel of the panel batholites.

Galicia has vast min the medicinal resources in itssubsoil, as there are more than three hundred sources stered, the ich twenty used by spas (Dirección Xeral de Industria, Enerxía e Minas, 2003). The use of the integration of the start of the start peutic means, dates back to Roman times. During the nite enth century the thermal

²⁵ peutic means, dates back to Roman times. During the nite of this century the thermal baths experienced golden age, with several spas, but at theend of this century a long





crisis began. resent more emphasis is being placed on their recreational and aestheticaspects than on their curative potential. Recently, Our ense was designated by the Galician Regional Parliament as the Thermal Capital of Galicia due to their therapeutic hot springs, representing the second highest position in thermal water of the Iberian Peninsula.

Most of these sources reach the highest water temperatures in Spain of about 71 °C. The abundance of these springs in Galicia is associated with the ithologic type and the soil fracturation.

The chemistry of thermal waters has attracted the attention of numerous studies to understand the processes that have influence on the recharge of underground water from its origin to estimate its resource importance and potential exploitation. In this way, some studies have been developed with the thermal waters of the area of Ourense city, the Thermal Capital of Galicia due to their therapeutic hot springs (González-Barreiro et al., 2009; Delgado-Outeiriño et al., 2009).

¹⁵ Carballiño is a municipality in the Spanish province of Ourense and has an area of 54 km². It has famous thermal spas together with multiple streams that bathe the countryside.

With the renewed interest in thermomineral waters, the principal aim of this study was to characterize the chemical equilibrium state of waters from Carballiño, as well as the thermodynamic conditions influencing water-rock interaction. As a secondary

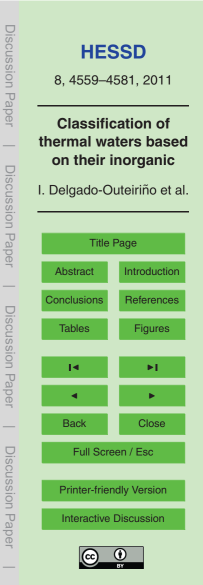
²⁰ as the thermodynamic conditions influencing water-rock interaction. As a secondary line of interest, this study aims to determine the temperature of the water within the reservoir.

2 Experimental

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2.1 Geochemical and hydrogeological setting

²⁵ From the geological point of view, Carballiño is situated on a granite crystalline substrate, which could be divided into two study areas, Northeast and Southwest (Fig. 1).



The North Eastern part consists of schist with granitic injections and two-mica granite rocks of adamellites, in which several intrusions of gneiss and schist are present. It is in this area where 10 of the 15 selected thermal waters (1-2-3-9-10-11-12-13-14-15) are located. Upwelling of these waters would occur in areas of contact between granite rocks and matematical (aphieta). This granitic material is alongly appaging the selected thermal waters (1-2-3-9-10-11-12-13-14-15) are located.

⁵ rocks and metamorphic materials (schists). This granitic material is closely associated with metamorphism and migamtization. The granite rocks of adamellites should be formed by anatexis in deeper areas, when the main Hercynian metamorphism reached the maximum temperature.

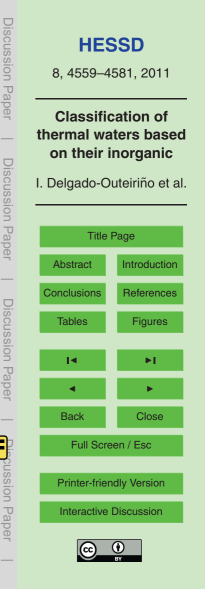
The most abundant rock material in the Southwest area is one of the most com-¹⁰ mon facies of granite, medium- to coarse-grained porphyritic granodiorite, which is composed of quartz, feldspar, plagioclase, muscovite and biotitic. Only one of the 15 thermal water here loca Beran Spa (8), with a temperature of 27 °C (Table 1). The rest of the selected thermal waters are related with the schistose material found in this area (4-5-6-7) with the highest temperatures (>40 °C).

The macro-fracturation of this area is represented by two large families of fractures that interact with each other. These families correspond to fracture N 20°W and N 130°W. An intense river network uses the lineation of these structures and it is observed along Miño river and also in its tributari circuit and provides the upwelling of springs.

20 **2.2** Sampling procedure in thermal water sites

In this study 15 thermal water samples were collected in April 2008 (Table 1).

Water samples were collected by immersing amber glass bottles at the points of emission. All samples were placed in a portable cooler, with ice, immediately after collection to prevent bio alteration; in the stored at 4°C until analysis within the next 24 h.



2.3 Chemical analysis

Temperature, pH and electrical conductivity of waters were measured in the field. Aluminium, boron, caesium, strontium and zinc were determined by Inductively Coupled Plasma Mass Spectrometry (Thermo X series II ICP-MS). Calcium, iron, magne-

sium and manganese were measured by atomic absorption spectrophotometry (Varian SpectrAA-250 Plus). Limit, sodium and potassium were analysed by atomic emission spectrophotometry rian SpectrAA-250 Plus Iloride, fluoride, sulphur, nitrite, nitrate, ammonium, phosphate and sulphate anions were determined by capillary electrophoresis (Thermo Capillary Electrophoresis using UV/Vis detector Cristal 100). Bi carbonate was determined by acidimetric titration. Analytical errors are generally <5% for the main components.

2.4 Thermal water chemistry

The combined use of $\frac{CI/SO_4, CI/HCO_3 \text{ and } (CI+SO_4)/HCO_3 \text{ and the Hill-Piper diagram}}{(Piper, 1944)}$ leaded to an improved classification of the the Hill-Piper diagram.

- The statistical methods used for data analysis of the samples vertices principal component analysis (PCA) and partial least squares regression (PLS-regression) (López-Chicano et al., 2001; Cruz and França, 2006; Cerón et al., 2009). The PC model was calculated on the auto scaled (namely, columns were mean-centred and scaled to unit variance) data. This was done to focus the analysis on in-between sample
 variations and unify the importance of each variable independently of the concentration levels. The model was further validated by cross-validation, visual inspection of loadings, argument interpretation to as certain the presence of a meaningful interpretation for are presented in the method of regression by PLS has been used extensively in
- chemometrics, where they have found a wide field of application. To attach a weight ing to each variable, the data obtained were divided by the standard deviation of each series and later processed by means of PLS2 algorithm of Unscrambler program, utilizing the method of "cross validation". "Leverage correction" validation method may



result in over-pristic validation results.Statistical analysis was carried out using the following statistical programmes: Unscrambler version 9.1 (Camo Process AS, 2004, www.camo.no) and Statgraphics version 5.1 both for Windows.

2.5 Geothermometers and hydrogeochemical modelling

Silica Geothermometers were used to obtain the most precise data possible about the theoretical reservoir temperature of our evetem and it was carried out using the thermodynamic database WATEQ4F.dat (the nd Nordstrom, 2001) included in the PHREEQC package (Parkhust et al., 1990). The PHREEQC package was also used for the geothermometric modelling.

10 3 Results and discussion

3.1 Chemical composition of waters and ionic ratios

The measurements taken on thermal waters are reported in Table 1.7 PH values range from 7.2 to 9.4, indicating alkaline nature thermal waters (Delgado-Outeiriño et al., 2009; López-Chicano et al., 2001). Anions are mostly represented by CO₃H⁻ (47–366 mg L⁻¹), followed by CO₃⁻² (n.d.–57 mg L⁻¹), Cl⁻ (9.3–29 mg L⁻¹), SO₄⁻²(4.0– 38 mg L⁻¹), F⁻ (n.d.–16 mg L⁻¹) and NO₃⁻ (n.d.–0.91 mg L⁻¹). Among the cations, Na⁺ is the main dissolved species (28–97 mg L⁻¹), followed mainly by Ca⁺² (2.2–16 mg L⁻¹) and K⁺ (0.60–6.6 mg L⁻¹). Temperature varies between 13° and 46°C. The electrical conductivity ranges from 137 to 629 µS cm⁻¹. A global analysis of the inorganic pattern of the selected waters derives in the distinction of only one type of waters as Fig. 2 shows: sodium-bicarbonated waters. Galicia was affected by the hercynianorogeny and these types of waters occur in the internal areas of post-orogenic fracture zones.

Moreover, PCA was carried out to reduce the structure of the data set to two dimensions. The total variance explained by these two components accounts for the 69%



(42% PC1 and 27% PC2) of the variability (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot of the selected samples shows two clusters (Fig. 3). The principal plan plot other, mainly by their higher Ca⁺², Mg⁺², Fe⁺², water random principal plan plot other, mainly by their higher Ca⁺², Mg⁺², Fe⁺², water hardness and height. The sequence Ca⁺² >Mg⁺² >Na⁺ is similar to the general depth sequence for groundwater composition outlined by Chebotarev (1955). When Ca⁺² >Mg⁺² >Na⁺ it means that young/surface water is present. A young fraction in a confined aquifer suggests possible modern recharge, continuity with the sequence samples that are distinguished due to their temperature, S⁻², SO⁻²₄, Si, *Li*, Na⁺ and dry residue contents. In this cluster, the springs belong to Prexiqueiro I, II, III, Cortegada and Laias.

Several ratios between different elements were also investigated and the most interesting results were observed between CI^{-}/SO_{4}^{-2} , CI^{-}/HCO_{3}^{-} and $(CI^{-}+SO_{4}^{-2})/HCO_{3}^{-}$ (Table 2). The CI^{-}/SO_{4}^{-2} ratio shows the interaction between water and rock (López-Chicano et al., 2001); a higher ratio value would indicate that this water has evolved over much longer time at depth and, therefore, it would interact with the rock. The same conclusion could be drawn with ference to the spatial evolution of the CI^{-}/HCO_{3}^{-} and

 $(Cl^- + SO_4^{-2})/HCO_3$ ratios (Lóppz-Chicano et al., 2001).

15

The elevated mobility of *Li* is related to temperature (Chan et al., 1994). It is found in high computations in thermal waters, and for this reason it is a good tracer for use in geochical investigations of hypermal systems (Prondi et al., 1973; Brondi et al., 1983). The concentration of *Li* in water depends a finite the water-rock contact time (Fidelibus and Tulipano, 1990) and, therefore, *Li* content could be used as an indicator of the residence time (Edmunds and Smedley, 2000; Sánchez-Martos et al.,

25 2004). Moreover, Leeman and Sisson (1996) found boron in very different geological environments, associated with the presence of volcanic rocks, geothermal processes, and with materials deposited in very saline environments. Because it is highly soluble, it tends to concentrate in environments that have a limited water circulation, like in evaporites or brines of marine or continental origin (Uhlman, 1991). Other authors

HESSD 8, 4559-4581, 2011 **Classification of** thermal waters based on their inorganic I. Delgado-Outeiriño et al. **Title Page** Introduction Abstract Conclusions References Tables **Figures** Back Close Full Screen / Esc **Printer-friendly Version** Interactive Discussion

Discussion Paper

Discussion Paper

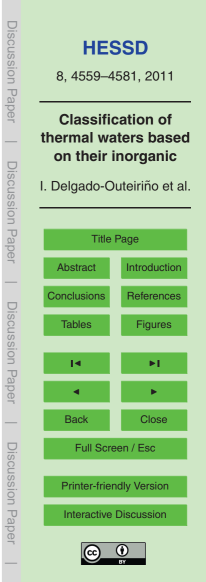
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consider that the elevated boron concentration in some connate waters is directly related to the content of K, *Li*, Mg, Sr and I (Macpherson and Land., 1). The high values recorded in thermal waters may be due to the alteration of volume rocks and hydrothermal activity (Risacher, 1984; Risacher and Firtz., 1991). In this respect the

- influence of temperature on its liberation has been noted (Arnorsson and Andresdottir., 1995). In order to find relationships between a set of the main compositional variables (variables X) and Li or B (Y variable) for data obtaining from the selected water samples, PLS-R was chosen. The selected algorithm was able to correlate a block of X variables with Li variable, giving a regression coefficient of 0.9988 for a model with two
- principal components and 0.8996 with *B* variable. Figure 4 shows a partial squares regression plots. The results obtained show clear separation, based on the first two principal components, between clusters I and II. In cluster I the samples 1, 2, 3, 8, 9, 10, 11, 13 and 14 with lower *Li* and *B* content were grouping. In cluster II the samples 4, 5, 6, 7 and 15 with higher *Li* and *B* content were found. The same clusters were found by PCA with the same distribution and therefore, the same results were found
- in both analysis. It could be proved that samples of cluster I were the youngest ones or they could be mixing continuity. Nevertheless in cluster II were the samples with longest water-rock contact time.

4 Geothermometer results

One of the most important applications of geochemistry for geothermal resources is using chemical geothermometers to give valuable information about what is happening in the reservoir. The accuracy of a geothermometer application is based on two assumptions. The basic assumption is that a temperature-dependent equilibrium is attained between fluid and minerals in the reservoir. It is further assumed that the composition of a fluid is not affected by chemical reactions in the upflow of geothermal system zones where cooling occurs (Wei, 2006). Various geothermometers have been developed to predict reservoir temperatures in geothermal system (Tole et al., 1993).



Geothermometers that have provided better results for these alkaline systems were those based on dissolved silica (SiO₂-chalcedony or SiO₂-quartz) and Na-K. These two techniques reflect the state of thermal equilibrium solution these systems with respect to quartz-chalcedony-albite, and potassium feldspar, respectively. In the present work, silica geothermometer was chosen due to the geochemical setting of Galicia.

Table 3 shows the equilibrium temperatures for quartz, chalcedony, kaolinite and k-mica. The chalcedony and or quartz equilibrium temperature also present for comparison. As can be in the calculated temperature of the quartz is the quartz chalcedony geothermometers are in the range from 70 to 100 °C and from 30 to 64 °C, respectively. It could be deduced from these results that quartz is the mineral phase, which rules the equilibrium state of the silica, as it was previously reported by Michard (1990), when he studied the behaviour of several elements in deep hot waters from granitic areas of Europe.

Two groups could be distinguished through the results obtained for both geother-¹⁵mometers. The first group would be integrated by the thermal waters, which reached the equilibrium at lowest temperatures (between 70–87 °C for quartz geothermometer and between 33–56 °C for chalcedony geothermometer) and can be shown in table 3 included the samples 1, 2, 3, 8, 9, 10, 13 and 14. The second group formed by the thermal waters 4, 5, 6, 7, 12 and that reached the equilibrium at highest temperature between 92–102 for quartz geothermometer and between 62–72 °C for chalcedony geothermometer).

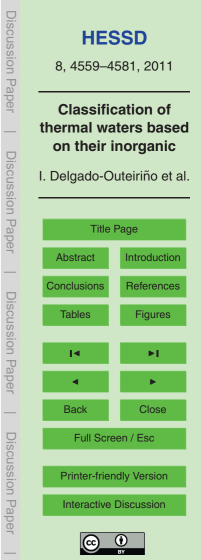
All of these results a with the previous obtained by PCA and PLSR, where also two groups were found. In group I the selected water samples could be in the previous with surface waters and therefore, the residence time in the reservoir and the previous of group II.

4.1 Hydrogeochemical modelling

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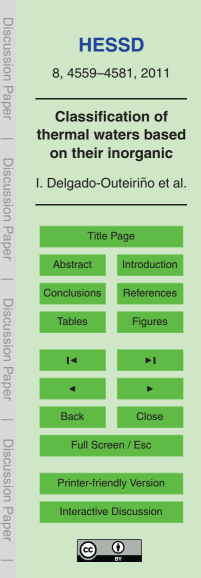
Geothermometers are based on the assumption that specific temperature-dependent mineral-solution equilibria are attained in the geothermal reservoir. Nevertheless it is



also advisable to study the fluid saturation equilibrium with the hydrothermal minerals in the reservoir (Reed and Spycher, 1984). In order to know the state of this equilibrium, saturation index (SI) was used and is the logarithm of the ratio (at each temperature) between the solubility product of a certain mineral by hydrolytical reaction (Q) and its

- ⁵ equilibrium constant (*K*). In this way, the inter-relationship between the lithologies encountered around the waters and their chemical composition could also be explained. All minerals in equilibrium at the same temperature converge to SI = 0; SI < 0 for an undersaturated solution, and SI > 0 for a supersaturated solution. In the present work, chalcedony, quartz, calcite, kaolinite and k-mica were selected to calculate the equi-
- ¹⁰ librium state for the selected thermal waters. Table 4 presents the SI for the selected minerals, calculated at the pH and temperature measured in the field. The studied waters are saturated with respect to quartz, k-mica, chalcedony and kaolinite, with the exception of Cortegada Baños and Partovia I. Other authors (López-Chicano et al., 2001) have been also observed super-saturation with respect to quartz in geothermal
- ¹⁵ fluids in Southern Spain. The lower saturation indices observed for chalcedony could be explained to its lower water solubility. The thermal waters 1, 2, 3, 5, 6, 8, 12, 13, 14 and 15 are under-saturated with respect to calcite whereas the samples 4, 7, 9, 10 and 11 and are super-saturated with respect to the same mineral, which is probably due to the cationic change of these thermal waters (D'Amore et al., 1987). Instead, the high est saturation index for kaolinite and K-mica could only be the result of a preferential
- circulation through feldspar and mica.

In Fig. 5 shows the variation of saturation indices SI with temperature for quartz and chalcedony kaolinite phases, considering a temperature interval between 20°C and 120°C; The intersection of SI curves at zero saturation indexes gives the equilibrium temperature (D'amore et al., 1987; Tole et al., 1993; López-Chicano et al., 2001). In this way, two groups could be also distinguished through the results obtained depending on the reservoir temperatures for the selected thermal waters. A group formed by the thermal waters, which reach the state of equilibrium at the highest temperatures, between 85 and 110°C (Fig. 5a) for quartz phases, and between 45 and 85°C (Fig. 5c)



for chalcedony and kaolinite phases. These thermal waters were the samples 4, 5, 6, 7, 12 and 15. The rest of the selected thermal waters would reach the state of equilibrium at the lowest temperatures between, $63^{\circ}C$ and $85^{\circ}C$ (Fig. 5b) for quartz phase, and between $35^{\circ}C$ and $65^{\circ}C$ (Fig. 5d) for chalcedony and kaolinite phases. SI obtained for these samples could depend on re-equilibrium processes during the ascent of the fluid to the surface.

5 Conclusions

5

In this paper the chemistry of major and trace inorganic elements in 15 thermal waters discharging in the council of Carballiño (province of Ourense) were present cussed. The results of the results o of bicarbonated waters the sodium. Graphical representations of CI/SO₄, CI/HCO₃ and $(CI+SO_4)/HCO_3$ ratios shows interactions between water and rock. These results were proved by PCAand PLSR, were the samples grouped in two clusters related with the water age and depth. Results of the geothermometric modelling as well as of geothermometers also agreed with the results obtained by the previous analyses and 15 analysis and also two groups of waters were detected. A group formed by thermal waters that reach the equilibrium at highest temperatures (between 85 and 110°C for IS for guartz and between 92 and 102 °C for guartz geothermometer, and between 45 and 85°C for IS chalcedony, and between 62 and 72°C for chalcedony geothermometer), which are under-saturated with respect to calcite. The second group of thermal waters 20 would reach the equilibrium at lowest temperatures (between 63 and 85 °C for IS quartz and 70 and 87 °C for quartz geothermometer, and between 35–65 °C for IS chalcedony and, between 33-56 °C for chalcedony geothermometer) and are super-saturated with respect to calcite. Comparable results were obtained for equilibrium temperatures ob-

tained modelling of the equilibrium states and by geothermometers with an error band of ± 10 °C, because of the equilibrium status at = h, chemical reactions at different temperatures (precipitation of kaolinite or calcite).



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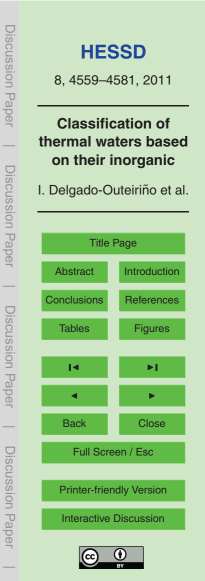
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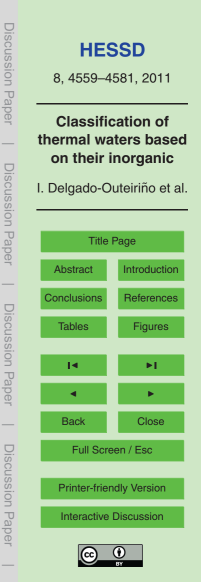


Table 1 samples		amp	olin	ıg	sit	tes,	and	d in	-sit	e a	Ind	lab	ora	tor	y m	nea	asu	ren	nen	ts o	of tl	he	15	the	rm	al v	vat	er		2
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Place (sample)		UTM (H29) Y	Height (m)	t pH	т (°С)	C (µS cm ⁻¹)	Na ⁺ mg L ⁻¹	<i>Li</i> ⁺ mg L ⁻¹	K ⁺ mg L ⁻¹	Ca ⁺² mg L ⁻¹	Mg ⁺² mg L ⁻¹	Sr ⁺² mg L ⁻¹	Fe ⁺² mg L ⁻¹	Zn ⁺² mg L ⁻¹	$^B_{mg L^{-1}}$	WH	CI ⁻ mg L ⁻¹	NH+4 mg L ⁻¹	F ⁻ mg L ⁻¹	CO ₃ mgL ⁻¹	Si mg L ⁻¹	S ⁻² mg L ⁻¹	CO ₃ H ⁻ mg L ⁻¹	Cs ⁺ mg L ⁻¹	Al ⁺³ mg L ⁻¹	SO ₄ ⁻² mg L ⁻¹	NO- mg L ⁻¹	DR		
A Rañoa (1) Ponterriza (2) Brues (3) Prexigueiro I (4)	569378	4696805 4697055 4699511 4678390	352 345 342 122	8.7 8.0 8.3 8.7	21 17 27 44	163 185 196 363	40 43 43 56	0.15 0.15 0.26 0.57	0.94 1.2 1.4 3.0	6.4 5.0 9.2 4.0	0.21 0.58 0.52 0.25	0.10 0.09 0.14 n.d	0.020 0.030 0.020 n.d	0.070 0.42 0.090 0.18	0.26 0.38 0.23 0.63	26 20 31 14	9.5 11 14 23	0.78 1.4 1.6 2.9	5.8 8.3 7.0	10 14 14 33	26 n.d. 33 54	0.23 n.d. 1.3 0.71	60 60 57 73	11 19 28 49	49 35 18 60	8.0 7.0 16 16	0.13 0.91 0.43 0.37	162 100 176 288	_	-

Harbara (1)	Impart 01 (1) Impart 01 (1)<	Impact 01	Hands (1)	Hands (1)	lace (sample)		UTM	Height	pН	T	C	Na ⁺	LI ⁺	K ⁺	Ca+2	Mg+2	Sr ⁺²	Fe ⁺²	Zn+2	B 1	WH	CI	NH+4	F-	CO31	Si	S-2	CO3H	Cs ⁺	AI+3	SO ₄ ⁻² mgL ⁻¹	NO
Interniza (1) 57874 4488730 5345 8.0 17 185 43 0.15 12 5.0 0.88 0.09 0.20 0.42 0.38 20 11 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 19 0.000 0.000 0.42 0.38 20 11 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 0.000 0.000 0.42 0.38 20 11 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 0.000 0.000 0.42 0.38 20 11 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 19 0.000 0.000 0.42 0.38 20 11 1.4 8.3 14 n.d. n.d. 60 19 9.35 7.0 0.91 19 0.000 0.000 0.42 0.38 20 11 1.4 8.3 14 n.d. n.d. 60 19 9.35 7.0 0.91 19 0.000 0.000 0.42 0.38 11 1.2 5.2 2.8 1.0 16 0.43 13 57 18 4.3 14 0.01 1.2 5.2 2.8 1.0 18 0.31 15 7.2 14 10 7.1 10 14 14 14 1.2 5.2 1.2 1.0 14 11 1.2 5.2 1.2 1.0 14 11 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2	Territar (2) 500776 469601 345 840 17 186 43 0.15 12 5.0 0.88 0.09 0.200 0.42 0.38 20 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 0.92 0.32 0.14 0.200 0.40 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 7.2 0.2 1.4 0.20 0.42 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 72 0.2 1.4 0.20 0.42 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 72 0.2 1.4 0.20 0.42 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 72 0.2 1.4 0.20 0.40 0.00 0.50 1.5 14 25 2.2 1.4 1.5 15 57 42 0.25 1.5 14 0.40 0.40 10 0.40 10 0.40 0.25 1.5 14 0.1 11 1.5 5.2 1.5 17 42 0.25 1.6 4 7.4 16 n.d. 1.4 0.1 0.4 0.25 1.5 14 0.25 0.41 11 1.5 5.2 1.5 17 42 0.25 1.5 14 0.41 8.3 14 0.1 0.8 0.40 10 0.40	Territar (2) 500776 469601 345 840 17 186 43 0.15 12 5.0 0.88 0.09 0.200 0.42 0.38 20 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 0.92 0.32 0.14 0.200 0.40 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 7.2 0.2 1.4 0.20 0.42 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 72 0.2 1.4 0.20 0.42 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 72 0.2 1.4 0.20 0.42 0.23 11 1 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 500076 469601 12 5.1 12 5.0 0.89 72 0.2 1.4 0.20 0.40 0.00 0.50 1.5 14 25 2.2 1.4 1.5 15 57 42 0.25 1.5 14 0.40 0.40 10 0.40 10 0.40 0.25 1.5 14 0.1 11 1.5 5.2 1.5 17 42 0.25 1.6 4 7.4 16 n.d. 1.4 0.1 0.4 0.25 1.5 14 0.25 0.41 11 1.5 5.2 1.5 17 42 0.25 1.5 14 0.41 8.3 14 0.1 0.8 0.40 10 0.40	Teneriza (1) Teneriza (2) Ten	Teneriza (2) 50007 4698705 345 8.0 17 185 43 0.15 12 5.0 0.88 0.09 0.030 0.42 0.38 20 11 14.8 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 magazine (1) 55802 4697305 122 8.4 47 358 56 0.56 1.3 4.0 0.25 1.4 0.00 0.09 0.42 0.33 11 1.4 8.3 14 n.d. n.d. 60 19 35 7.0 0.91 magazine (1) 55802 4697305 122 8.4 47 358 56 0.56 1.3 4.0 0.25 1.4 0.00 0.99 0.691 14 25 2.2 n.d 53 34 0.7 17 4.6 0.4 19 0.5 18 0.47 magazine (1) 55802 4697305 122 8.3 40 456 73 11 12 8.4 47 358 56 0.56 1.3 4.0 0.25 1.4 0.00 0.99 0.691 14 25 2.5 n.d 6 1.3 54 0.7 17 4.6 0.4 19 0.5 18 0.47 magazine (1) 55802 4697305 122 8.3 40 456 73 11 1 1.3 5 3.2 0.56 n.d. 0.4 0.02 0.59 0.25 1.5 14 25 1.5 17 42 0.25 116 0.47 10 0.4 10 0.4 0.25 1.5 14 0.5 18 0.4 10 0.4 10 0.4 0.0 0.4 0.0 0.25 1.5 14 0.5 18 0.4 10 0.4 0.0 0.4 0.0 0.4 0.0 0.4 0.0 0.4 0.0 0.5 0.5 1.4 0.0 0.0 0.4 0.0 0.5 0.5 1.4 0.0 0.0 0.4 0.0 0.0 0.4 0.0 0.5 1.5 14 0.5 18 0.4 1 0.5 0 0.4 0.0 0.5 0.5 1.5 14 0.5 0.4 0.0 0.5 0.5 1.5 14 0.5 0.4 0.4 0.5 0.5 0.4 0.0 0.5 0.5 0.5 0.4 0.0 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5			(H29) Y	(m)		(.)															mgL							mg L ⁻¹
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Discussion Paper

Discussion Paper

	CI^{-}/SO_4^{-2}	CI ⁻ /HCO ₃	$(Cl^{-}+SO_{4}^{-2})/HCO_{3}^{-}$
A Rañoa 📃	3.2	0.27	0.36
Ponterriza 1	4.3	0.32	0.39
Brues 🏹	2.3	0.42	0.59
Prexigueiro I	3.9	0.54	0.68
Prexigueiro II	1.8	0.38	0.60
Prexigueiro III	2.3	0.40	0.61
Cortegada Baños	4.8	0.42	0.51
Beran Balneario	2.3	0.31	0.44
Partovia I	2.8	0.38	0.52
Partovia II	2.9	0.33	0.44
Partovia III	2.8	0.35	0.47
Gran Balneario	2.7	0.36	0.49
Arcos Carballiño	3.6	0.38	0.49
O Xardín	2.7	0.31	0.43
Laias Balneario	3.2	0.32	0.45

Table 2. Some ionic ratios (in $meq L^{-1}$) of interest in the selected thermal waters.

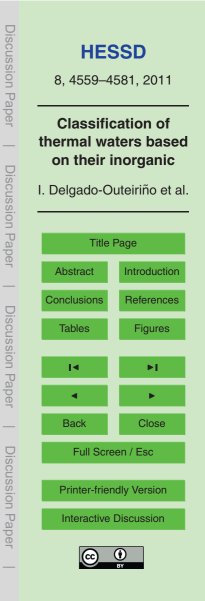


Table 3. Quartz, chalcedony, kaolinite and k-mika equilibrium temperatures (°C) using the ther-
modynamic database WATEQ4F.dat (Ball and Nordstrom, 2001). Equilibrium temperatures for
the geothermometers of SiQ_2 -quartz and chalcedony are also shown.

Places (sample)	Chalcedony	Quartz	Kaolinite	K-mica	T- quartz	T- chalcedony	Group
A Rañoa (1)	41	73	41		73	41	I
Ponterriza (2)	58	73	60	97	80	49	I
Brues (3)	52	82			83	52	I
Prexegueiro I (4)	65	101	56	82	95	65	II
Prexigueiro II (5)	67	97	58	85	92	62	II
Prexigueiro III (6)	70	103	55	94	95	62	11
Cortegada (7)	70	82	70	72	92	64	II
Beran (8)	40	70	63	81	72	40	I.
Partovia I (9)	42	63	35	65	70	48	I
Partovia II (10)	45	75	40	70	72	50	I.
Partovia III (11)	35	64	35	64	75	40	I.
Gran Balneario (12)	73	101	45	78	98	71	II
Arcos (13)	54	85	45	75	87	56	I
O Xardín (14)	40	65	65	73	65	33	I.
Laias(15)	75	100	85	110	102	72	II



Discussion Paper

Discussion Paper

Discussion Paper

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Chalcedony Quartz Calcite Kaolinite k-mica	A Rañoa 1 0.220 0.660 -1.04 8.94 15.7	Ponterriza 2 0.340 0.800 -2.57 3.58 6.4	Brues 3 0.260 0.680 -1.48 8.85 15.2	Prexigueiro I 4 0.320 0.710 0.220 0.640 4.60	Prexigueiro II 5 0.650 0.760 -0.240 2.27 6.40	Prexigueiro III 6 0.240 0.620 -0.350 0.850 4.56	Cortegada Baños 7 0.0300 0.420 0.510 -0.440 3.31	Beran 8 0.0800 0.510 -0.600 2.91 6.43	Partovia I 9 -0.100 0.290 -0.380 2.92	Partovia II 10 0.0600 0.470 0.230 0.630 4.22	Partovia III 11 0.110 0.540 0.0600 1.01 4.63	Gran Balneario 12 0.400 0.820 -0.110 1.46 5.51	0.340 0.800 -0.130 2.20	O Xardín 14 0.240 0.710 -1.48 3.42 6.10	Laias 15 0.250 0.620 -0.530 3.45 7.36	—	Abstract Conclusions Tables	Introduction References Figures
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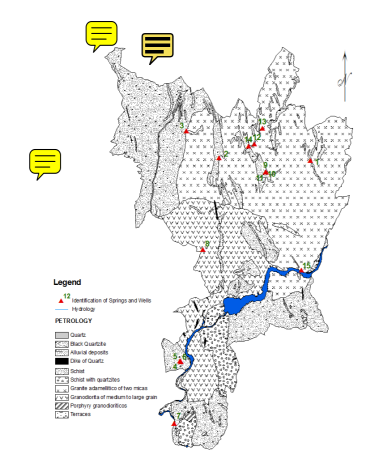
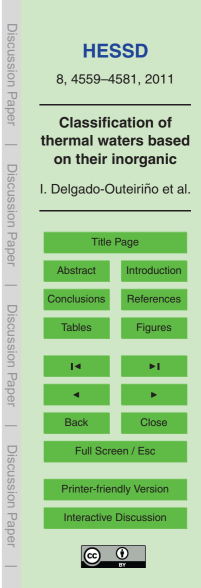
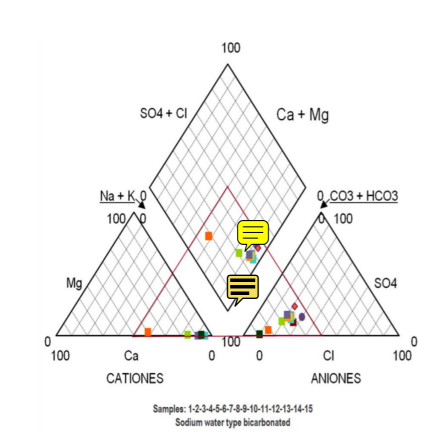
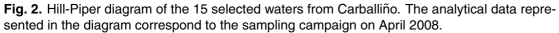
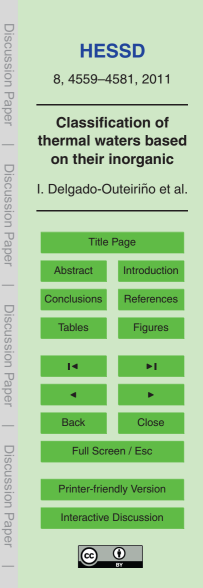


Fig. 1. Hydrogeological map with the sampling sites.









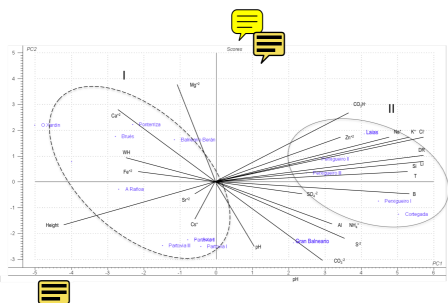
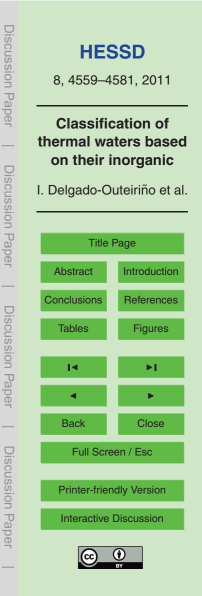


Fig. 3. Prinicpal component analysis on the selected thermal waters. Cluster I, samples with the shortest water-rock contact time, lowest equilibrium temperatures and under-saturated with respect to calcite. Cluster II, samples with longest water-rock contact time, highest equilibrium temperature and super-saturated with respect to calcite.



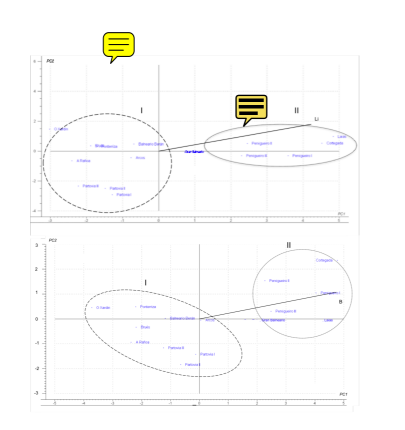
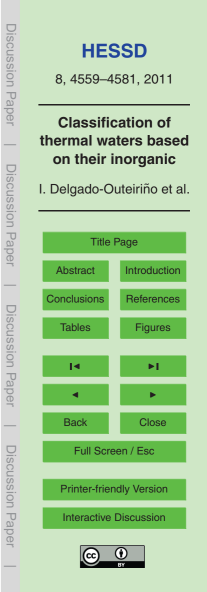


Fig. 4. Partial Least-squares regression plots for the selected samples, showing the *X* loading weights and the *Y* loadings for *Li* and *B*. Regression coefficient of 0.9988 and 0.8996 for *Li* and *B*. Two clusters are also distinguished depending on *Li* and *B* contents.



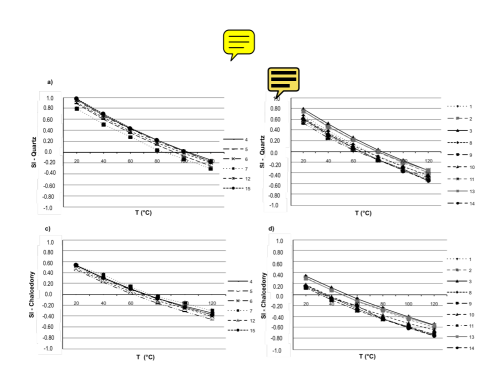


Fig. 5. Result of the geothermometric simulations: (a) Quartz saturation index for the samples with equilibrium temperature between 85-110 °C, (b) Quartz saturation index for the samples with equilibrium temperatures between 63-85 °C, (c) Chalcedony saturation index for the samples with equilibrium temperatures between 45-85 °C, (d) Chalcedony saturation index for the samples with equilibrium temperatures between 35-65 °C.

