Are There Benefits from Thermal Bacteria for Health? The Hydrogenome Role

Patrick Pascoal-Ferreira 1,2,3, Daniel Glez-Peña 4,5, Carla Miranda 3,6, Patricia Poeta 3,6, João Coutinho 7, Florentino Fdez-Riverola 4,5, Ana Torrado-Agrasar 8,9, María Luisa Rúa 8,9 and Gilberto Igrejas 1,2,3,*

1. Introduction

Water is considered the basis for biodiversity on our planet, it is part of all life processes and it has a primordial function in the existence of life as we know it [1]. The properties of...
water give it fundamental characteristics for the structural organization and for the three-dimensional assembly of proteins and nucleic acids [2], acting also in the accomplishment of the diverse biological functions of these molecules [3].

For thousands of years, humans have lived and seized numerous lands, grown and developed due to this invaluable resource that is water [4]. However, over time, humans began to observe the beneficial effects that some waters had on health, these waters became known as the thermal waters and some civilizations even attributed to them a strong religious and sacred character [5–7]. Both the ancient Greeks and the Romans surmised the beneficial properties of the thermal waters well and used them to alleviate joint pain, muscle pain or the treatment of skin diseases [8,9]. Thermal waters are all-natural mineral waters with therapeutic applications or health beneficial effects, emerging from the inside of a thermal spring, independently of the source temperature, according to Medical Hydrology [10]. According to the European Directive 2009/54/EC, natural mineral waters are defined as microbiologically wholesome and originated from an underground water table or deposit, emerging from a natural spring or bore. These waters are characterized by nature (mineral content and trace elements, among others) and original purity. In addition, these water characteristics can confer beneficial effects to health, considering the assessment of the main following parameters, such as physicochemical, microbiological (absence of pathogenic microorganisms and parasites), hydrological and geological [11].

Due to the scientific advances, the therapeutic experience of physicians and the new studies of hydrology, pharmacology and biochemistry, thermal waters have once again taken on great importance [9] in the prevention and treatment of some heart diseases [12], skin diseases [13–15], respiratory diseases [16,17] or rheumatic diseases, such as osteoarthritis, fibromyalgia, ankylosing spondylitis, rheumatoid arthritis and chronic back pain [18–20].

The specific therapeutic purposes of each thermal water are closely related to the microbial diversity and the different physical and chemical characteristics they present, such as pH, temperature, total mineralization, salts concentration and the nature of the ionic components, among others [7,10,11,21]. Based on the chemical composition, the natural mineral waters are classified as sulphated, bicarbonated, chlorinated, sulphurous, gasocarbonic and hiposaline waters. Each water chemical composition is associated with a specific therapeutic indication. For instance, sulphurous waters are indicated for reumathic and musculoskeletal, dermatologic, gynecologic and respiratory diseases; gasocarbonic waters are indicated for circulatory and digestive systems; bicarbonate thermal waters are indicated for respiratory, urinary, digestive and endocrine diseases [10,21]. These characteristics depend on their respective lithostructural and tectonic context or geographic location [7,21]. Water, when reacting with the various components present in the rocks, obtains some constituents that will modify its composition. A recent study reported that silica-rich thermal waters showed dermatologic properties with clinical benefits in case of atopic dermatitis and psoriasis, reducing the macrophages and keratinocytes proliferation and cell metabolism [15]. Additionally, the presence of microbial communities associated with thermal waters also contribute to the diversity and specialization of the hydromineral richness, according to some authors [7,13,21]. However, thermal waters with the geographic location in seismic areas should be monitored carefully, since the earthquake can show a high impact on water safety and stability, including the restoration of microbial composition [22].

There are a number of studies, where the importance of microbial communities in thermal waters has been described in obtaining new molecules of biotechnological interest [23–26]. However, only recently a few authors [13,21,27,28] have verified that the microbial composition of these waters may also play a greater role in the health benefits. Thermal waters have unique microbiomes related to the physicochemical characteristics previously described; however, the presence of some active prebiotics may stimulate the growth of bacterial species beneficial to health, especially if they are already present in its natural microbial content [13,27]. For instance, the antirheumatic properties of
sulphurous-bromine-iodine thermal water have been associated with the presence of the followed bacteria belonging to genera Geothermobacterium, Thermus, Syntrophomonas, Desulfomonile, Thiofaba and Thermodesulfovibrio, according to Paduano et al. [21]. Moreover, these characteristics can contain or also allow the growth and colonization of pathogenic microorganisms like Legionella. The predominant genera identified in water samples obtained from the borehole of an Australian balneotherapy centre were Sphingobium, Agrobacterium and Thiobacillus [29].

Given the benefits presented directly related to health, there is a need to understand and study the total content and diversity of microorganisms present in this water type. With the development of the next generation sequencing (NGS) and the increase of computational capacity in bioinformatics, it is possible to have access to the total hydrogenome present in the thermal and natural mineral waters [22,26,30]. Therefore, the term hydrogenome comprises all the genomes of the microorganisms present in a given aquatic environment. This study of hydrogenome may be used as an auxiliary tool in the practice of medical hydrology, since this new approach makes possible the analysis of the composition of the taxonomic and functional profiles of the microorganisms existing in the thermal waters, including the raw material of the thermalism or balneotherapy, and increasing the safety in the used water. The use of NGS and bioinformatics tools has the main objective of assisting classical microbiology methods by allowing the direct and rapid study of the genome of the microbial communities from the most diverse environmental samples, including the identification of uncultured microorganisms [26,30].

The Iberian Peninsula is rich in hot springs with therapeutic purposes. In particular, the region of Galicia in Spain and the northern region in Portugal have a strong thermal tradition through the exploration of these resources [31–34]. For these reasons, the aim of this pilot study was to characterize the hydrogenome of five thermal waters of these two regions of the Iberian Peninsula, using NGS and the molecular technique of the 16S rRNA gene.

2. Materials and Methods

2.1. Sampling Site

This study was carried out in two regions of the Iberian Peninsula, namely in the region of Galicia (Spain) and in the northern region of Portugal.

The region of Galicia is essentially formed by a vast outcrop of metamorphic rocks (crystalline rocks) and igneous (granitic rocks). In this Spanish province, there is also a great variety of hot springs, with more than three hundred sources, of which 20 are used by spas [33]. The chemical composition of the waters emanating from this region is largely conditioned by several factors, such as atmospheric gases, aerosols, surface erosion of rocks and soils (including their biological constituents), the dissolution and precipitation of minerals from the surface and the existence of a certain “thermal anomaly” in the region [33]. The chemical composition of the Galician waters is mainly bicarbonated sodium and calcium waters, resulting from the occurrence of carbonation reactions and hydrolysis of the minerals present in the subsoil, among them feldspar [35,36]. Galician waters have as main therapeutic indications at rheumatological, respiratory, dermatological, digestive and hepatobiliary levels [37].

Portugal considered one of the richest thermal countries of the European Union, comprises more than fifty thermal centers [15]. One of the main Portuguese regions to natural thermal mineral waters is located in the Transmontano-Duriense, which presents a huge variety of hot springs [38], typologies and occurrences. These waters are mainly of meteoric origin, that is to say, they result from a deep infiltration coming from rainwater [39]. The thermal waters currently in operation in this region may have been infiltrated for about 10,000 years, at the beginning of the Holocene, placing the hydrothermal richness of the Trás-os-Montes region on a plateau of great national prominence [39]. The natural mineral waters of the Trás-os-Montes region are mainly bicarbonate, sodium, gasocarbon, fluoride and acid waters [31,40]. These waters are still geared to therapeutic practice at the level of
the cardiovascular, digestive, respiratory, metabolic-endocrine, dermatological, rheumatic and musculoskeletal systems and nervous system [15,31,40,41].

2.2. Sample Collection

Water samples (50L) were collected from November 2017 to March 2018, in the following points, ES_TI (42.346583, −7.877306), ES_PR (42.254466, −8.167167) and ES_BUR (42.334706, −7.864560), which were related to the samples collected in the Galicia region in Spain, while the points PT_SA (41.855243, −7.185342) and PT_CH (41.738313, −7.473070) corresponded to the samples collected in the northern region of Portugal. The collection was performed after flushing water for 1 min, filling the bottles up to the top and closing them with caps. During the process of water collecting in the respective sources, the measurement of its emergency temperature was carried out using a digital thermometer (Table 1).

Table 1. Physicochemical composition of the analyzed thermal waters.

| Physicochemical Parameters | Units | Water Samples |
|----------------------------|-------|---------------|
|                            |       | ES_BUR | ES_PR | ES_TI | PT_CH | PT_SA |
| Condutivity (C)            | µS    | 956    | 461   | 447   | 2320  | 1455  |
| Dry residue (DR) at 110 °C | mg/L  | 656    | 374   | 365   | 1626  | 998   |
| pH                         |       | 7.5    | 7.3   | 6.8   | 6.8   | 6.1   |
| Emergency temperature      | °C    | 64     | 50.4  | 40    | 70    | 15    |
| Lithium (Li)               | mg/L  | 0.98   | 0.74  | 0.54  | 2.33  | 1.84  |
| Sodium (Na)                | mg/L  | 222.8  | 82    | 87    | 703.7 | 205   |
| Potassium (K)              | mg/L  | 9.26   | 4.67  | 3.82  | 65.32 | 23.9  |
| Rubidium (Rb)              | mg/L  | 0.14   | 0.04  | 0.03  | 0.6   | 0.12  |
| Cesium (Cs)                | mg/L  | 0.18   | 0.11  | 0.06  | nd    | 0.23  |
| Calcium (Ca)               | mg/L  | 8.90   | 3.1   | 3.78  | 20.46 | 72.8  |
| Magnesium (Mg)             | mg/L  | 0.64   | 0.12  | 0.16  | 4.54  | 37.1  |
| Strontium (Sr)             | mg/L  | <0.01  | 0.04  | 0.03  | 0.37  | 1.41  |
| Iron (Fe)                  | mg/L  | 0.02   | 0.02  | <0.01 | 0.23  | 0.35  |
| Boron (B)                  | mg/L  | 0.70   | 0.85  | 0.51  | 0.79  | 0.52  |
| Aluminium (Al)             | mg/L  | <0.01  | <0.01 | <0.01 | <0.01 | <0.01 |
| Manganese (Mn)             | mg/L  | 0.04   | <0.01 | <0.01 | 0.02  | 0.24  |
| Lead (Pb)                  | µg/L  | 0.30   | <0.2  | <0.2  | <0.2  | <0.2  |
| Chromium (Cr)              | µg/L  | <1     | <1    | <1    | <1    | <1    |
| Cadmium (Cd)               | µg/L  | <0.5   | <0.5  | <0.5  | <0.5  | <0.5  |
| Copper (Cu)                | µg/L  | 1.18   | <1    | <1    | <1    | <1    |
| Zinc (Zn)                  | µg/L  | 10.17  | 1.6   | 0.9   | <1    | 2.2   |
| Total Arsenic              | µg/L  | 6      | 14    | 13    | 79.20 | <1    |
| Arsenic III                | µg/L  | nd     | 13    | 10    | 20.30 | nd    |
| Arsenic V                  | µg/L  | nd     | 1.79  | 3.25  | 58.90 | nd    |
| Cobalt (Co)                | µg/L  | <1     | <1    | <1    | <1    | <1    |
| Molybdenum (Mo)            | µg/L  | 1.17   | 3.5   | 1.1   | 0.74  | <0.2  |
| Vanadium (V)               | µg/L  | 20.00  | <1    | 20.00 | <1    | <1    |
| Chloride (Cl−)             | mg/L  | 28.44  | 28.83 | 15.86 | 38.30 | 10.26 |
| Ammonical nitrogen (NH4+)  | mg/L  | 0.43   | 0.01  | 0.5   | 1.62  | 0.35  |
| Sulphite (SO3−)            | mg/L  | <1     | 4.3   | <1    | 1     | <1    |
| Sulphate (SO4−)            | mg/L  | 3.5    | 29.93 | 17.76 | 11.29 | 1.26  |
| Fluoride (F−)              | mg/L  | 16.63  | 21.97 | 14.76 | 8.01  | 0.54  |
| Sulphide (S2−)             | mg/L  | <0.05  | 9.05  | 3.38  | <0.05 | 0.46  |
| Carbonate (CO3−)           | mg/L  | 0      | 12    | 12    | 0     | 0     |
| Bicarbonate (HCO3−)         | mg/L  | 585.6  | 122   | 195.2 | 1769  | 1207.8|
| Nitre (NO2−)               | mg/L  | 0.64   | <0.05 | <0.05 | <0.05 | <0.05 |
| Nitrate (NO3−)             | mg/L  | <0.05  | <0.05 | <0.05 | 2.62  | <0.05 |
| Phosphate (PO4−)           | mg/L  | 1.53   | <0.05 | <0.05 | <0.05 | <0.05 |
| Iodide (I)                 | µg/L  | 3.58   | 1.9   | 1.3   | nd    | 1.3   |

nd: not determined.
2.3. Physicochemical Analysis

All the water samples were analyzed for the following physicochemical parameters: micronutrients (iron, copper, zinc, manganese, boron, cobalt, molybdenum, vanadium, iodine, fluorine and arsenic), cations (sodium, potassium, rubidium, calcium, cesium, magnesium, strontium, lead, cadmium, aluminum and ammoniacal nitrogen) and anions (chlorine, sulphites, sulphates, sulphides, carbonates, bicarbonates, nitrates, nitrates and phosphates). These analyses were carried out in the Laboratory of Soils and Plants Joaquim Quelhas dos Santos at the University of Trás-os-Montes and Alto Douro and at the Center for Scientific and Technological Support for Research of the University of Vigo, using standard methods. The values are shown in Table 1.

Taking into account the physicochemical parameters, the thermal waters collected in two regions of the Iberian Peninsula were classified chemically using the Piper diagram, indicating their similarities according to the cations and anions (expressed in mEq/L) represented by two different triangles [42,43]. Additionally, the principal component (PC) analysis and cluster analysis were performed, in order to evaluate the similarities and differences of the physicochemical composition of waters. The PC analysis was performed using the Excel XLStat statistical package (version 2.11.1, Addinsoft, Paris, France) [44]. The experimental data on the physicochemical characterization of the waters (Table 1) were first normalized, from these a correlation matrix was obtained, the own values of each main component and the percentage of the variance was explained for each main component. As a criterion for the choice of PCs, the number of components explaining most of the variability (greater than 70% of the total variance) was considered [45].

2.4. DNA Extraction and 16S Amplicon Sequencing

Water samples were filtered through 0.22 µm filters (Sartorius Stedim Biotech, Gottingen, Germany), in order to isolate the microorganisms, present in the thermal water. Total genomic DNA extraction from samples was performed using the DNeasy® PowerWater® kit (Qiagen, Hilden, Germany), following the manufacturer’s instructions. After extraction of DNA, amplification and preparation of libraries were performed according to the 16S metagenomic sequencing library preparation (Part # 15044223, see A, Illumina Inc., San Diego, CA, USA). The amplicons were obtained by the amplification of the V3-V4 regions of the 16S rRNA gene of both Bacteria and Archaea, using the following primers 341F (5' - CCTACGGGNGGCWGCAG-3') and 785R (5' - GACTACHVGGGTATCTAATCC-3') [46], and specific adapters of the Illumina MiSeq platform (Illumina Inc., San Diego, CA, USA) [47–49]. The libraries obtained were quantified and validated through a Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA). After obtaining the libraries, sequencing was performed through the Illumina MiSeq platform, using the MiSeq Reagent Kit v3, according to the manufacturer’s protocol and in a 2 × 300 bp paired-end system. All of the sequencing was performed at the Madrid Science Park.

2.5. Bioinformatic Analysis

The obtained sequencing data were analyzed using bioinformatics tools, such as FASTQC(Babraham Bioinformatics, Cambridge, UK) [50] and Metagenomic Rapid Annotations using Subsystems Technology (MG-RAST server, http://www.mg-rast.org (accessed on 12 September 2018)) [51]. The FASTQC tool was used to analyze the quality of the FASTQ files from the MiSeq sequencer (16S rRNA sequencing). This tool allows for the reporting of a wide variety of information related to the quality profile of the readings obtained, and for evaluating the guanine–cytosine (GC) content and the abundance of adapters [52], which may still be present after the initial preprocessing carried out by the company Madrid Science Park.

This MG-RAST server is a bioinformatic tool that allows the functional and taxonomic annotation of sequences of environmental samples (metagenomics) through the comparison with databases such as KEGG, M5NR, Genbank, PATRIC, RefSeq, SwissProt and SEED, among others. This tool also offers quality control, annotation and various methods for
analyzing different types of data, including phylogenetic and metabolic reconstructions, and the ability to compare more than one metagenome [51].

Reads were taxonomically annotated by similarity searching against M5NR database, respectively, with default parameters (maximum e-value cutoff of $10^{-5}$, minimum identity cutoff of 60% and minimum alignment length cutoff of 15).

3. Results and Discussion
3.1. Physicochemical Classification of Natural Thermal Mineral Waters

The five water samples analyzed in this study showed distinct physicochemical and microbiota composition. The physicochemical analysis of water samples showed that the pH values vary between 6.1 and 7.5, thus indicating slightly acidic, neutral and alkaline waters (Table 1). Regarding the temperatures recorded in the various thermal waters, these were found to vary between 15 and 70 °C. Normally, higher temperatures favor the appearance of more mineralized water [7], however, water PT_SA has very high mineralization values even though it is a water with a low emergency temperature (15 °C), which may be related to the fact that emergency temperature is often much lower than the maximum temperature reached in the reservoir, as it decreases along the ascending hydraulic circuit [33,53].

According to European legislation, water with a ≤50 mg/L dry residue is classified as hyposaline or very little mineralized, 50–500 mg/L is low mineralized, 500–1500 mg/L is classified as mineralized water and water with dry residue above 1500 mg/L is classified as hypersaline, presenting strong mineralization [54]. Thus, water samples ES_PR and ES_TI can be classified as low mineralized waters, ES_BUR and PT_SA as mineralized waters and PT_CH as being hypersaline water with values of mineralization above 1500 mg/L (Table 1). These results are confirmed by the values of electrical conductivity. Since this parameter allows us to evaluate its degree of mineralization as results from the relationship between the content of dissolved minerals in the water and the resistance that it offers the passage of electric current.

Based on the Piper diagram, thermal waters of the two regions of the Iberian Peninsula were classified chemically (Figure 1), the Piper diagram analysis showed that there are mainly three water groups, two of them in a transient phase (Figure 1). The first group included the water samples ES_BUR, ES_TI and PT_CH, classifying them as bicarbonated sodium waters. In addition, these samples presented high values of sodium (Na) and potassium (K), but also of bicarbonate anions (HCO$_3^-$) (Table 1). The second group, the sample ES_PR is located in one of the transition zones, between the bicarbonated sodium waters and the sulfated or chlorinated sodium waters. The classification can be justified when analyzing these results together with Table 1, since the sample ES_PR had the highest values of sulfate anions (SO$_4^{2-}$) and sulfite (SO$_3^{2-}$), compared to the other samples. Regarding the third group, the water sample PT_SA is also located in a transition zone, between the sodium bicarbonated waters and the bicarbonated calcium or magnesian waters. Additionally, the sample PT_SA had the highest values of magnesium (Mg) and calcium (Ca), but also had a high content of sodium (Na) and bicarbonate anions (HCO$_3^-$), thus placing it in the transition between the bicarbonated sodic waters and the bicarbonated calcium or magnesian waters.

The analysis of principal components based on the all analyzed physicochemical parameters (Table 1) showed that considering the first two main components (PC1 and PC2), the obtained results can explain 71% of the total system information (Table 2). However, the addition of a third major component (PC3) can explain 95% of the cumulative variance. Therefore, only the first three main components were selected for the next analysis.
Figure 1. Classification of the thermal waters analyzed in this study, based on their anions and cations composition, using the Piper diagram.

Table 2. Obtained values of the correlation matrix of the thermal waters analyzed, for the four principal components (PCs).

|       | PC1    | PC2    | PC3    | PC4    |
|-------|--------|--------|--------|--------|
| Own value | 13.82  | 8.92   | 7.54   | 1.73   |
| Variability (%) | 43.19  | 27.87  | 23.55  | 5.393  |
| Cumulative variance (%) | 43.19  | 71.05  | 94.61  | 100.00 |

The projection of the first two main components (PC1 and PC2) and the main components (PC1 and PC3) are demonstrated in Figure 2, respectively. Regardless of the PC1 and PC2 components or the PC1 and PC3 components can be observed that the five samples of thermal waters in both projections were always grouped in four groups, although a displacement occurs in the positioning of the waters in the quadrants. Considering this reason and that the PC1 and PC2 components present more than 70% of the variance in comparison to 66.74% for the PC1 and PC3; the thermal waters were clustered into the following groups: (1) ES_BUR, (2) PT_CH, (3) PT_SA and (4) ES_PR and ES_TI.

Figure 2. Projection of the first two main components (PC1 and PC2), showing 71.05% of cumulative variance (A) and projection of the main components (PC1 and PC3), showing 66.74% of cumulative variance (B).

The results of the PC analysis represented graphically using the “biplot” function with the scale of species and localities (Figure 3), allowed visualizing the variables that most
influence the classification of the thermal waters. The variables with greater weight in each component are represented (square cosine $> 0.5$).

![Figure 3](image)

**Figure 3.** Results of the PC analysis of the main components PC1 and PC2 represented graphically, using the “biplot” function with 71.05% of cumulative variance.

Considering the water samples into four groups, among which ES_PR and ES_TI formed an independent cluster located in the opposite (and therefore negatively correlated) quadrant relative to water PT_CH. In turn, the PC analysis positions the water ES_BUR in the quadrant opposite PT_SA. The water samples ES_PR and ES_TI demonstrated a negative correlation with the PC1 and PC2 components, and the ions that allowed them to be grouped were mainly composed of sulfur ($\text{SO}_3^{2-}$ and $\text{S}_2^{2-}$) and $\text{CO}_3^{2-}$. This fact was confirmed by physicochemical results, in which these two waters showed the richest in sulfur compounds (Table 1). This type of water exerts a number of beneficial effects on health, namely anti-inflammatory, keratoplastic and antipruritic properties, also having antibacterial and antifungal properties due to the presence of sulfur [10,55]. The sulfur may also interact with oxygen radicals present in the deepest layers of the epidermis, generating sulfur and sulfur dioxide, which can be transformed into pentaionic acid ($\text{H}_2\text{S}_5\text{O}_6$), acid responsible for the antibacterial and antifungal activity of sulfur water [55].

In relation to the other waters, the variables that most contribute to their grouping in different groups are the cations (Mg, Ca and Sr) for the water sample PT_SA and the nitrogen compounds ($\text{NO}_3^-$ and $\text{NH}_4^+$), conductivity, dry residue, Na and Rb for the sample PT_CH. For water ES_BUR, the pH is the only variable that contributes to classify it, considering the components PC1 and PC2.

### 3.2. Characterization of Microbiota Communities of Natural Thermal Mineral Waters

A total of 804,158 sequence reads were obtained from the five water samples by the sequencing of the 16S rRNA gene, with an average length of 426 base pairs (bp) (Table 3). About 99.95% of the reads submitted from the thermal water samples were maintained after the quality control step of the server itself. In addition, evaluation of the GC content demonstrated good global GC contents ($> 50\%$) in all water samples. These sequences were deposited in the NCBI Short Read Archive (GenBank accession numbers SRX5099959-SRX5099963, [https://www.ncbi.nlm.nih.gov/bioproject/?term=PRJNA508462](https://www.ncbi.nlm.nih.gov/bioproject/?term=PRJNA508462) (accessed on 20 May 2021)). According to Browne et al. [56], the optimal coverage for GCs content should be range between 45 and 65% in the metagenome dataset. While GC content outside of this interval can show coverage biases, indicating difficulty on genome sequencing.
For each analyzed water sample, the rarefaction curve was calculated through the MG-RAST server. These rarefaction curves represent the richness of the different species annotated as a function of the number of reads presented, demonstrated that practically all samples tend to approach an asymptote before 100,000 reads, that is, most species present in the samples have already been discovered (Figure 4). Taking into account that all samples reached a stable plateau, obtaining more than 100,000 reads after the quality control of the MG-RAST server (Table 3) and a larger number of reads would not increase the count of species obtained [21,57]. These results demonstrated a reliable sequencing of the microbiota community and there is still a great diversity of unknown microorganisms.

Table 3. General metagenomic annotation data on the MG-RAST server.

| Samples | Number of Reads before Quality Control | Number of Reads after Quality Control (%) | Average Reads Length (bp) | % of GC Content after Quality Control |
|---------|--------------------------------------|------------------------------------------|---------------------------|--------------------------------------|
| ES_BUR  | 164,464                              | 164,377 (99.95)                          | 421                       | 58 ± 5                               |
| ES_PR   | 157,364                              | 157,320 (99.97)                          | 435                       | 54 ± 5                               |
| ES_TI   | 189,567                              | 189,439 (99.93)                          | 428                       | 53 ± 5                               |
| PT_CH   | 175,886                              | 175,741 (99.92)                          | 415                       | 59 ± 4                               |
| PT_SA   | 116,877                              | 116,838 (99.97)                          | 430                       | 52 ± 5                               |
| Total   | 804,158                              | 803,715 (99.95)                          | 426                       |                                      |

GC: Guanine–cytosine.

Figure 4. Rarefaction curves of the bacterial 16S rRNA gene sequences for each water sample obtained from the MG-RAST server. The total number of sequence reads analysed is plotted against the number of species observed in the same community.

Regarding the number of species, the samples tended to cluster into two distinct groups, ES_BUR, PT_SA and PT_CH formed a first group, and ES_PR and ES_TI formed a second group (Figure 4). This may be related to the different temperatures recorded in their waters. The temperature is one of the factors that most affects the microbial growth, as these increases accelerate the enzymatic reactions, and the growth is faster [58]. This process occurs until reaching a maximum temperature from which some of the cellular components can begin to denature irreversibly (proteins, nucleic acids, etc.), thus stopping the microbial growth [58,59]. Therefore, each microorganism has its own optimum growth temperature in which the enzymatic reactions occur at the maximum possible speed. Paduano et al. [21] reported the highest bacterial diversity and complexity of the thermal waters in the lowest temperatures, varying between 36 and 44 °C. In the case of mesophilic microorganisms, their temperatures can vary between 15 and 60 °C [23]. Taking into account that in this study, the water samples of group two (ES_TI and ES_PR) had temperatures between 40 and 50.4 °C, this could be one of the main factors that contributed to the appearance of
a higher species richness (number of species) compared to group one. Group one included the water samples (ES_BUR, PT_CH and PT_SA) with more extreme temperatures, which probably could be related to a lesser amount of microbial diversity. In addition, the extreme environments present numerous challenges for the microorganisms that inhabit them, challenges that may require an adaptation of all cellular machinery for both psychrophilic microorganisms and for thermophilic and hyperthermophilic microorganisms [60]. Thus, given that both low and high temperatures affect the structure and function of microbial cells [23,61]. Into group two is expected that the existence of microorganisms in these waters is strongly conditioned by the type of adaptations they have, showing a lower degree of species richness compared to the samples of another group (Figure 4). Some studies demonstrated that few microbial communities are adapted to conditions of high (>60 °C) or low temperatures (<20 °C), resulting, therefore, in a low diversity compared to mesophilic environments [62,63]. For instance, Everroad et al. [59] reported that the only Aquificales members, in particular Sulfaurogenumium, grew above 67 °C from a slightly alkaline sulfide-containing hot spring in Japan. In addition, these authors observed that with the decrease of temperature, the primary productivity increased through the anoxygenic and oxygenic phototrophs, and consequently, the community diversification.

Comparison of Microbial Taxonomic Profiles

The taxonomic classification based on 16S rRNA gene sequences analysis allowed determining the composition of the microbial communities associated to the different waters analyzed in the two regions of the Iberian Peninsula, through the MG-RAST server, according to Ribosomal Database Project (RDP) classification by interactive graphs (Krona) created to each sample (Figures 5 and 6). Krona graphs represent an important metagenomic visualization tool, facilitating the study and access to bioinformatic data [64]. In this study, the taxonomic profile showed that between 99.0% and 99.9% of assigned reads were from the Bacteria realm and the remainder from Archaea.

Figure 5. Analysis of the taxonomic distribution of the microbial community in the Spanish thermal waters, based on the 16S rRNA gene sequences.
The water sample ES_BUR showed that 99.95% of the sequences belong to the realm of Bacteria, of which 60% were of the Proteobacteria (45% Betaproteobacteria, 14% Gammaproteobacteria and 1% Deltaproteobacteria), 8% Deinococci, 6% Nitrospira, 4% Actinobacteria, 2% Aquificae and 4% belonged to minority classes. However, 16% of these bacterial sequences were not classified (Figure 5).

The sample ES_PR, 99.99% of its sequences belonging to the Bacteria kingdom, of these sequences, 22% belong to the phylum of Proteobacteria (9% Epsilonproteobacteria, 7% Betaproteobacteria and the remaining were distributed among the other classes, such as Deltaproteobacteria), 13% belong to the Nitrospira class, 10% to the Firmicutes class (3% Bacilli and 7% Clostridia class), 2% to the Actinobacteria class, 1% to the phylum Verrucomicrobia and Bacteroidetes, and the remaining 5% belong to minority classes. While 46% were unclassified bacterial sequences.

The sample ES_TI had 99.99% of obtained sequences belonging to the Bacteria kingdom, of which 50% were unclassified, 22% belong to the phylum of Proteobacteria (15% Gammaproteobacteria, 5% Deltaproteobacteria and the remaining 2% were distributed among the other classes, such as Betaproteobacteria), 17% belongs to the Nitrospira class, 5% to the Firmicutes class (3% Bacilli class and 2% Clostridia), 2% Bacteroidetes, 1% to the Actinobacteria classes and the remaining 3% belong to the minority classes.

In the water sample PT_CH, the major of sequences were associated to the realm of Bacteria (99.95%), of which 42% belongs to the phylum Proteobacteria (27% Betaproteobacteria, 13% Gammaproteobacteria, 1% Deltaproteobacteria and the remaining 1% is distributed among the other classes, as is the case of Alphaproteobacteria), 18% to the Deinococcaceae class, 16% to the Aquificae class, 14% were not classified and the remaining 10% belong to the minority classes, such as Actinobacteria and Nitrospira classes (Figure 6).

In relation to the sample PT_SA (Figure 6), 99% of the sequences belongs to the realm of Bacteria, of these sequences, 78% were of the Proteobacteria phylum (74% Epsilonproteobacteria, 3% Betaproteobacteria and the remaining 1% was distributed by other classes, such as Gammaproteobacteria), 19% are unclassified and the remaining 2% are distributed by the minority phyla, such as the phylum Firmicutes. In the case of this water sample, in particular, it is verified that 1% of the total sequences obtained were classified as belonging to the realm of Archaea, of these sequences, 100% are of the Euryarchaeota phylum.

In all water samples of this study, the phylum Proteobacteria (22% to 78%) was the most abundant, following the phylum Deinococcus-Thermus (8% to 18%), Nitrospira (5% to 17%), Aquificae (2% to 16%) and Firmicutes (2% to 10%). In addition, the Proteobacteria phylum was clearly the most representative phyla in the following water samples PT_CH, ES_BUR and PT_SA, compressing the class Betaproteobacteria (ES_BUR and PT_CH), Epsilonproteobacteria (PT_SA and ES_PR) and the class Gammaproteobacteria (ES_BUR, ES_TI and PT_CH). The phyla and classes with a representation ≥10% are shown in Figure 7.
Figure 7. Comparison of the most representative bacterial diversity communities at the phylum and class level (≥10% of the sequences) of the analyzed natural thermal mineral water samples in this study, including the unclassified bacterial sequences (US).

Proteobacteria have relevant biological importance since this phylum is mostly constituted by Gram-negative bacteria and these have a high medical, veterinary, industrial and agricultural interest [65,66]. In general, most Proteobacteria are mesophilic bacteria, that is, with optimal growth temperatures between 15 and 60 °C, however, there are some bacteria that may be thermophilic, as in the case of *Thiomonas thermosulfata* or psychrophiles, such as *Polaromonas* [65]. The possibility of these bacteria being present in both psychrophilic, mesophilic and thermophilic environments confirms our results obtained in Figure 7, where there is also the existence of Proteobacteria in water with temperatures between 15 and 70 °C.

Some studies have demonstrated that Proteobacteria represent an important component of the skin microbiota [13,40,67,68]. However, although this phylum has already been detected in several metagenomic studies, the microbiota of cutaneous Proteobacteria still remains little described [69]. The microbial communities of the human skin have strong capacities to interact with their environment and can thus establish a relationship between skin and environmental ecotypes in the same species of Proteobacteria [70]. Considering the various therapeutical properties associated with natural thermal mineral waters, such as psoriasis, acne, alopecia, eczema and (chronic) atopic dermatitis, among others [55]; some studies have demonstrated that the microbial composition of these waters may also play a greater role in the health benefits [21,27,28,71]. The One Health concept recognizes that human health is strongly linked to animal health and the environment [69].

In relation to the Firmicutes phylum in this study, the samples that presented a greater amount of sequences were ES_PR and ES_TI (Figure 5). This phylum has some interesting characteristics for scientific research, in particular by including some genera of aerobic or anaerobic bacteria that produce endospores. These endospores are responsible for conferring resistance to conditions of high temperatures and other environmental stresses [72]. The Firmicutes phylum also presents genera such as *Staphylococcus, Enterococcus, Streptococcus, Clostridium, Bacillus* and *Lactobacillus*, being these genera are quite used in biotechnological processes. The appearance of microorganisms of the Firmicutes phylum in thermal waters with temperatures around 50 °C, has been frequently reported worldwide [73–75].

The species belonging to the Deinococcus-Thermus phylum show a great biochemical, physiological and phenotypic diversity [76], being these bacteria able to survive high doses of ionizing radiation, vacuum, high temperatures, desiccation, hydrogen peroxide and several other agents capable of cause DNA damage [77,78]. The presence of microorganisms of this phylum in the samples of water with higher temperatures (PT_CH and ES_BUR) corroborates the obtained results by several authors [78,79].
The microorganisms belonging to the Aquificae phylum are Gram-negative, do not form spores and are strictly thermophilic with an ideal growth usually occurring above 65 °C [80,81]. This optimum growth temperature corroborates our results, in which the Aquificae phylum is mostly present in the water sample PT_CH (with a temperature of 70 °C). Regarding their metabolism, most of the microorganisms belonging to the Aquificae phylum are bacteria that can oxidize hydrogen, but alternatively, thiosulfate or sulfur can also be used as energy sources [80]. The thermal stability found in many of the enzymes belonging to the genus Aquifex and other thermophilic bacteria of the Aquificae phylum, are of particular interest in industrial and biotechnological applications [80,81].

The Nitrospira Phylum is composed of bacteria capable of oxidizing nitrite, being the most abundant and diverse group of bacteria that act on nitrification [82]. Some studies have demonstrated its wide distribution in natural habitats, in soils [83], oceans [84] and hot springs [85]. Members of the genus Nitrospira belong to the most interesting microorganisms for biotechnology [86].

4. Conclusions

This study demonstrated that the thermal waters of the Galician region of Spain and the northern region of Portugal could be clustered into four distinct groups based on their physical–chemical composition. The waters ES_PR and ES_TI were high content in the sulfur compounds (SO\textsubscript{3}\textsuperscript{2−} and S\textsuperscript{2−}) and CO\textsubscript{3}\textsuperscript{2−}. For the sample ES_BUR, the variable that most contributed to its grouping was the pH. The sample PT_SA was rich in the cations (Mg, Ca and Sr) and the sample PT_CH was constituted by the nitrogen compounds (NO\textsubscript{3}− and NH\textsubscript{4}+), conductivity, dry residue, Na and Rb.

Based on the sequencing of the 16S rRNA gene, the water samples revealed a high diversity and a bacterial predominance of the Proteobacteria phylum (in water samples PT_CH, ES_BUR and PT_SA), followed by the phylum Firmicutes (in waters ES_PR and ES_TI), Deinococcus-Thermus (in waters PT_CH and ES_BUR), Aquificae (in waters ES_BUR and PT_CH) and Nitrospira (in waters PT_CH, ES_TI, ES_PR and ES_BUR); the abundance of Archaea was higher in water PT_SA (1%).

This innovative work allows for the first time to study the taxonomic diversity of microbial communities in the waters of the northern region of Portugal, and it also contributes to increasing the existing knowledge of the microbial diversity of the thermal waters of the Galician region. Although it was possible to identify most of the microbial composition of the studied waters used for therapeutic purposes; there are samples (ES_PR and ES_TI) that showed more than 40% of the obtained sequences as unclassified, still demonstrating an enormous gap in the knowledge of the microbial communities of these waters.

The study of these microorganisms that exist in these unique habitats was able to occur mainly due to the metagenomics and the new technologies of sequencing (NGS), allowing it to accede to the most diverse microbiomes. This knowledge has the possibility of leading to the discovery of new drugs, namely antibiotics, immunosuppressants, enzymes and new compounds of great importance for both medicine and biotechnology. Additionally, this work can contribute to increasing the safe use of these unique natural thermal mineral waters. The continuity of the work is essential, for instance analyzing the metabolic pathways of the microorganisms present in the waters and identifying how these metabolic pathways may be related to the diversity and specialization of the hydromineral richness. In addition, the elaboration of hydrogen OTEC will be allowed to group all hydrogenomes of the natural thermal minerals waters of the Iberian Peninsula. Finally, correlate the physical–chemical composition of the waters with the diverse microbial groups found in these habitats is essential to amplify the knowledge of these thermal waters.

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