Microbiological study of the dripping waters in Altamira cave (Santillana del Mar, Spain)

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The culturahle microbial populations in dripping waters from Altamira cave were studicd and compared with those of the ceiling rock. Water cornmunities have low proportions of gram-positive bacteria, and are mainly composed of gram-negative rods and caeci *(Enterobacteriaceae* **and** *Vibrionaceae),* **while those of ceiling rocks are mainly** *Streptomyces* **spp. The community differences are probably related to environmental cave conditions: high humidity, relatively low and stable temperature, water pH clase to neutrality and nature of the organic mattef. AH these factors seem to favor coloruzation and long-term growth of actinornycetes over other heterotrophic bacteria on ceiling rocks. rights reserved.**

Keywords: **dripping water;** *Enterobacteriaceae; Vibrionaceae; Streptomyces;* **crystal formation**

1. Introduction

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The Altamira cave, situated on the Cantabrian comice, Santillana del Mar, Spain, is known as the Sistine Chapel of Quatemary Art. The cave has the most important prehistoric paintings of Spain, and probably of the world, particularly the Polychromes Hall, which contains the majority of the Magdalenian paintings, about 14000 years old (Valladas et al., 1992).

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The cave, in a small calcareous hill, 158 m above sea level, was discovered in 1879. Since then, Altamira has suffered a series of changes related to its structure and the increasing growth in number of visitors, in such a way that the Polychromes Hall

was reduced to a small artificial chamber, wilh enviromnental conditions very different from the natural ones and leading to deterioration of the paintings.

Deterioration of paintings and microbial growth was related to the high number of visitors, which reached a daily flow of 1500 persons in the 1960s, increasing rapidly in the 1970s to 3000 daily, despite the first alarming signs of deterioration observed in the paintings. In 1976 a Commission in charge of conservation proposed the dosure of the cave, but this did not happen, for various reasons, until 1979. Since 1982 visits to the cave were reduced to 45 persons per day, but the conservation problems of the paintings still remain (Hoyos, 1993).

The nature and distribution of dripping waters in Altamira have been the basis of controversy. Villar et al. (1983) selected ten dripping water points·in the ceiling of the Polychromes Hall and observed that the monthly average of dripping water is 6 ± 1 l. **2. Material and methods** However, Hoyos (1993) stated that the dripping water flux is not constant, nor are its geochemical 2.1. *Sampling and sample location* characteristics.

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A preliminary study of bacterial communities in A sampling campaign was carried out to investithe dripping waters in Altamira cave was carried out gate the microbial population in the dripping waters by Somavilla et al. (1978). *Bacillus* and *Pseudo*- of the cave of Altamira at the end of January 1997. *monas* appeared as the most abundant genera with Gonzalez de los Reves-Gavilan et al. (1984) stated six and five species, respectively, followed by that the number of bacteria changed seasonally with *Flavobacterium* and *Erwinia* with two species. Five a peak in February. sampling points were located outside the Poly-
Dripping water from five different points along the chromes Hall which showed higher variability (ten cave (Fig. 1) were collected in sterile tubes (tripgenera) than two sampling points inside the Hall licate) and kept at 4° C until microbiological analysis with only three species of the genera *Bacillus* and was carried out two days after sampling. The points *Erwinia*. Uruburu et al. (1981) estimated cfu are located in the Kitchen Hall (EN-1), the gallery to (colony-forming units) of bacteria in two samples of the Polychromes Hall (EN-2 and 3), the Hall of the dripping water but no identification of genera and Walls (GM) and the Big Hall (GS) in a transect of species was provided. Hardisson et al., 1982 found about 800 m. A chemical analysis of the waters is differences in the cfu of bacteria between two shown in Table 1. samplings in November 1981 and February 1982. For comparison, two samples (P-5 and P-6) from Gonzalez de los Reyes-Gavilan et al. (1984) reported the surface of the Polychromes Hall ceiling and one that the dripping waters contained a considerable (C-EG) from the ceiling of the gallery conducting to number of bacteria, which was not eliminated this Hall (about 2 meters from EN-3) were studied. through rock filtration. These points were sampled by plate impressions

Recently, Groth et al. (1999) reviewed the growth from the surface. of actinomycetes in caves and hypogean environments. Two caves with rock art, Altamira and Tito Bustillo, were selected as a case-study. Within about 2.2. *Enumeration* 350 actinomycetes were identified by morphological, physiological and chemotaxonomic methods in Alta- Petrifilm aerobic count plates were inoculated with mira cave. Most of the actinomycetes growing on the 1 ml water sample. The plates contained a readysurface of ceiling and wall rocks were colonies from made medium for enumerating total aerobic bacteria 1 to 10 mm diameter, visible with the naked eye. Many isolates corresponded to strains obtained directly from the colonies. The genera *Streptomyces*, *Nocardioides*, *Amycolatopsis*, *Brevibacterium*, *Nocardia*, *Rhodococcus*, *Aureobacterium*, and the family *Micrococcaceae* were well represented. In Tito Bustillo cave, the surface of the rock was colonized by a large number of small, yellow, round colonies of about 1–2 mm. Direct isolates from the colonies were found to be *Streptomyces xanthophaeus* strains.

In this paper, the microbial population in the dripping waters of the Altamira cave is studied and compared with those of the ceiling and wall rocks using culture methods. The sampling value of the state of the Fig. 1. Altamira cave and dripping water sampling points.

Table 1 Chemical and microbiological analysis of dripping waters

Sample	pH	Т°С	CO ₂ ^a	CO, H^-	SO_4^{2-}	Cl^-	Ca^{2+}	$M\varrho^{2+}$	$Na+$	K^+	CFU m l^{-1}
$EN-1$	7.84	13.4	17.5	307.9	38.7	12.0	88.9	12.30	8.96	2.08	210
$EN-2$	7.62	13.7	25.0	457.3	5.0	9.0	122.5	16.45	7.39	0.41	TNTC ^b
$EN-3$	7.63	14.6	17.5	405.5	53.1	38.5	121.0	18.6	21.87	9.73	75
GM	7.63	14.2	25.0	350.6	51.7	21.5	116.6	7.97	13.28	2.87	310
GS	7.62	13.5	12.5	350.6	26.3	11.5	114.3	4.08	9.29	. . 50	TNTC

^a mg l⁻¹ of dissolved CO₂, all other anions and cations expressed in mg l⁻¹.
^b Too numerous to count.

the medium contains (per litre) 5 g tryptone, 2.5 g bacterial strain was expressed as percentage of total yeast extract and 1 g glucose, a cold-water-soluble activity of an isolate: $G_{(i)}$ %, and calculated accord-
gelling agent, and a tetrazolium indicator dye which ing to Kölbel-Boelke et al. (1988) from the number gelling agent, and a tetrazolium indicator dye which facilitate colony enumeration. Counts shown in of positive and negative characters of each isolate. Table 1 represent the mean on at least 6 Petrifilm Physiological activities of the community for digesplates for each triplicate sampling point. The plates tion of specific substrates was obtained from the were incubated at 28°C for 48 h. Some water number of isolates with positive and negative characsamples were inoculated directly in Sigma tryptone-
ters. soy agar (TSA) plates.

For comparison, samples from the rock surface 2.5. *Precipitation of salts* from the Polychromes Hall and galleries were enumerated on Rodac plates prepared with different All isolates were tested for salt precipitation using culture media: TSA, malt-yeast extract -MEY- (Laiz, B-4 medium composed of: 2.5 g calcium acetate, 4 g 1991), starch-casein -SC- (Küster and Willians, yeast extract, 10 g glucose, 15 g agar in 1 l distilled 1964), glycerol-asparagin -GA-, humic acid-agar water, pH 8 (Boquet et al., 1973). Precipitation was -HA- (Hayakawa and Nonomura, 1987), and also tested in liquid medium for the isolates with cycloheximide-agar -CA- (Dietz and Thayer, 1980). higher precipitation capacity in order to collect the The plates were incubated at 28° C for 48 h before crystals for further analysis. SEM and EDX, X-ray enumeration and were recounted after 8 weeks to diffraction and FT-IR were routinely used for this record slowly growing microorganisms. purpose.

2.3. *Identification*

After enumeration, individual colonies were randomly isolated and purified by streak plating on TSA 3.1. *Cell counts* until pure cultures were obtained. Isolates were characterized by morphological and physiological Petrifilm plates have been widely used for the properties, the latter using standard microbiological enumeration of contaminated liquids and foods methods, API and Biolog. Identifications were per- (Ginn et al., 1984; Matner et al., 1990; Byrne and formed with APILab Plus and Biolog databases. Bishop, 1991) and were considered a good choice for

pounds were calculated as in vitro activities from ever, in the sampling period dissolved carbon was

populations. According to the manufacturer (3M), API test reactions. Total physiological activity of one

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3. Results and discussion

this study. Table 1 shows Petrifilm counts at the 2.4. *Physiological activities* different sampling points. Dissolved-organic-carbon content in Altamira samples showed high variability The activities of decomposition of organic com-
depending on the dripping points and seasons. Howless than 5 mg C 1^{-1} (van Grieken, personal com-
The opposite was found in the samples from the munication). The concentration of organic materials ceiling of the Polychromes Hall and galleries. In the in recharge water and in aquifers is low, typically ceiling of the Polychromes Hall, the proportions of around 1 mg C 1^{-1} , a concentration believed too low gram-positive bacteria were threefold higher (95.7% to support to support life (Leenheer et al., 1974). In Altamira for P-5, 87.8% for P-6) near the paintings, or in the dripping waters, cell counts varied from 75 cfu ml⁻¹ access gallery to Polychromes Hall (84.6% for Cto concentration of colonies too numerous to count. EG). was reported by Stetzenbach et al. (1986) for a lated was *Aeromonas*, occurring in fresh waters and for 39 drinking water wells from which water was gallery and at the Big Hall sampling point. *A*.

contained low proportions of gram-positive (27.3%) the cave. *Acinetobacter* spp. comprised 54% of the relative to gram-negative (72.7%) bacteria (Fig. 2). total number of isolates from deep-well ground water It has also been shown that water communities (Stetzenbach et al., 1986). It was suggested that contain low proportions of gram-positive bacteria stimulation of growth of *Acinetobacter* sp. by low when compared with sediments or soils (Kölbel- concetrations of carbon reflects the ability of this Boelke et al., 1988; Wilson et al., 1983). organisms to effectively utilize a variety of carbon

Very close sampling points also showed a high Among the gram-negative bacteria isolated from variability. A similar variability (30–690 cfu ml⁻¹) the dripping waters, the most abundant genus isocontinuously working deep groundwater well. Wol-
ters and Schwartz (1956) obtained 10–32 cfu ml⁻¹ species most frequently isolated both at the entrance pumped from depths of 40–50 m. Kölbel-Boelke et *salmonicida* and *A. sobria* were the other al. (1988) found less than 100 cfu ml⁻¹, and in many *Aeromonas* species identified in samples obtained cases even less than 5 from a sandy aquifer from 5 m below the surface. were identified to very good levels of confidence, but the *A*. *sobria* strain did not utilize citrate (80% 3.2. *Bacteria identification* utilization for the species).

Acinetobacter spp. were also isolated from the Culturable isolates from Altamira water samples entrance and from the sampling points farther into sources and may in part explain its predominance in well water.

> Quantitatively important was also the genus *Enterobacter*, widely distributed in nature and occurring in fresh water, sewage and animal and human feces, represented by the species *Enterobacter amnigenus*. This bacterium was present in the dripping waters from the gallery leading to the Polychromes Hall (EN-3). Other bacteria identified in this sampling point were *Serratia liquefaciens*, *Chromobacterium violaceum*, *Pseudomonas fluorescens*, *Pseudomonas aeruginosa*, *Janthinobacterium lividum*, *Chryseomonas luteola*, *Xanthomonas maltophilia*, *Flavimonas oryzihabitans*, and *Kingella kingae*. Four gram-negative bacteria could not be identified.

The culturable gram-positive bacteria were represented by the genus *Bacillus*. The species *B*. *cereus*, Fig. 2. Distribution of bacteria in Altamira samples. Dripping B. circulans, B. subtilis, B. stearothermophilus, and water samples are summarized under WATER, P-5 and P-6 are B. polymyxa were mainly isolated from dripping samples from the ceiling of Polychromes Hall and C-EG from the water taken at the sampling points located at the cave access gallery to Polychromes Hall. entrance (Table 3). A microbial community based

Table 2 Gram-negative isolates of dripping waters of Altamira cave

Group	Identification	No isolates	O rigin ^a	Percentage of identification ^b
Facultatively anaerobic	Enterobacter amnigenus		$EN-3$	91.5
gram-negative rods	Serratia liquefaciens		$EN-3$	81.3
	Erwinia sp.		GМ	85.8
	Aeromonas hydrophila	3	GS, EN-1, EN-2	99.6
	Aeromonas sobria		$EN-1$	74.8
	Aeromonas salmonicida		$EN-3$	99.5
	Chromobacterium violaceum		$EN-3$	99.8
Gram-negative	Janthinobacterium lividum		$EN-3$	77.0°
aerobic/microaerophilic	Pseudomonas fluorescens		$EN-3$	97.3
rods and cocci	Pseudomonas aeruginosa		$EN-3$	97.0
	Chryseomonas luteola		$EN-3$	99.8
	Xanthomonas maltophilia		$EN-3$	88.0°
	Flavimonas oryzihabitans		$EN-3$	81.3
	Acinetobacter sp.		GS , $EN-1$	95.0
	Kingella kingae		$EN-3$	84.0°
	NI^a		EN-1, EN-2, GS	

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a EN: entrance of the cave, GM: Hall of the Walls, GS: Big Hall, NI: not identified.

^b An estimate of how closely the profile corresponds to the taxon relative to all the other taxa in the Apilab Plus data base.

c Identified with Biolog.

in rock art paintings from Atlanterra shelter (Gon- were obtained (56% with clear aerial mycelia). zalez et al., 1999). Seven isolates were identified as *Streptomyces* spp.

ping waters agrees with the observation of Kölbel- were *Erwinia amylovora*, *Chryseomonas luteola*, Boelke et al. (1988). They found very few ac- *Pseudomonas fluorescens* and *Bacillus sphaericus*.

twenty three isolates (Fig. 2) six were *Streptomyces rishiriensis*, *S*. *flavogriseus*, *S*. *xanthophaeus* and spp., and four gram-positive non-endospore produc- *Streptomyces* sp. were identified in addition to ing, calcium carbonate crystal precipiting strains *Flavimonas oryzihabitans* and *Xanthomonas* sp. (Fig. 3). These data clearly demonstrate that dripping water

almost exclusively on *Bacillus* spp. was also found P-6 yielded 27 cfu cm⁻², from which 41 isolates The absence of culturable actinomycetes in drip- and one as *Nocardioides* sp. Other bacteria found tinomycetes in 60 water and sediment samples. Four *Streptomyces* spp. precipitated crystals in B-4
Bacterial enumeration of a sample from ceiling medium (Fig. 3). Thirteen isolates were obtained Bacterial enumeration of a sample from ceiling medium (Fig. 3). Thirteen isolates were obtained rock surface (P-5) showed 30 cfu cm⁻². From from C-EG sample, among which *Streptomyces*

a EN: entrance of the cave, GM: Hall of the Walls, NI: not identified.

^b An estimate of how closely the profile corresponds to the taxon relative to all the other taxa in the Apilab Plus data base.

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microbial communities were different from those of rock.

3.3. *Physiological activities*

Dripping water isolates were tested for their ability to degrade different substrates. Table 4 shows total physiological activity of the dripping water isolates. Isolates from the families *Enterobacteriaceae* and *Vibrionaceae*, distributed homogeneously in all samples, showed the higher values, followed by some *Bacillus* species. Physiological activities of the isolates for digestion of specific substrates is shown in Fig. 3. Distribution of bacteria forming calcium carbonate. Expla-

Table 5, expressed as percentage frequency of posi-

nation as in Fig. 2. Histograms represent number of calcium

tive characters. This table compares the nation as in Fig. 2. Histograms represent number of calcium
carbonate-producing gram-positive and gram-negative bacteria
with respect to cfu.
by the isolates of the water community. All gram-

Table 4

Physiological activity of dripping water isolates			

 a^{a} G(i)%: percentage of total activity of a bacterial strain.

^b NI: not identified.

Table 5 Comparison of the 10 substrates digested with highest frequencies by isolates from dripping waters

Gram-negative isolates		Gram-positive isolates			
Substrate ^a	Isolates using substrate $(\%$ of total)	Substrate ^a	Isolates using substrate $(\%$ of total)		
ADH	54.2	GLU	100.0		
GLU	50.0	FRU	100.0		
ARA	41.7	MAL	100.0		
CIT	41.7	ESC	100.0		
GEL	41.7	CEL	88.9		
SAC	37.5	AMD	77.8		
MEL	29.2	GLYG	77.8		
AMY	25.0	GEL	77.8		
RHA	20.8	RIB	66.7		
INO	8.3	MNE	66.7		

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a ADH: arginine, CELlobiose, GELatine, MALtose, RIBose; AMD: starch, CITrate, GLUcose, MELibiose, SACcharose, AMYgdalin, ESCulin, GLYcoGen, MaNnosE, ARAbinose, FRUctose, INOsitol, RHAmnose.

maltose and esculin, however, gram-negative isolates showed comparatively lower frequencies for glucose either from water or rock samples was *Bacillus*, with (50%), arabinose (41.7%), saccharose (37.5%), and *B*. *cereus* in water and *B*. *sphaericus* in rock as melibiose (29.3%). In general, gram-positive bacteria isolates. Considering the total number of isolates, the presented higher positive reactions for carbon and percentage was much higher in the P-6 rock sample nitrogen sources than gram-negative bacteria, with than in dripping water and other rock samples. In the exception of rhamnose, lysine, ornithine and fact, the total percentage of crystal-producing bacarginine. The data demonstrate that endospore-form- teria was 18.2% for dripping waters, 43.9% for P-6, ing gram-positive rods degraded monosaccharides 17.4 for P-5 and 15.4% for C-EG. with the highest frequencies, but higher frequency of Canaveras et al. (1999) suggested that in Altamira occurrence of tested physiological activities were cave, some calcium and magnesium crystal deposits usually found among facultatively anaerobic gram-
originated by the action of bacteria. The bacteria negative rods. **isolated either from dripping waters or rock showed**

formations of calcite, aragonite, and hydromagnesite can be found (Canaveras et al., 1999). The presence calcium carbonate occurred on hyphae suspended of aragonite and hydromagnesite are highly sug- from the stalactite wall. He reported that the hyphae gestive of a microbial-mediated precipitation. To this function both as crystallization nuclei and as attachend, crystal formation with all isolated bacteria from ment, without which individual crystals would be dripping waters and ceiling rock was tested (Fig. 3). eliminated by the falling drop. The bacteria producing crystals in a B-4 medium were *Acinetobacter* sp., *Serratia liquefaciens*, 3.5. *Crystal analysis Chryseomonas luteola*, *Xanthomonas maltophilia*, *Flavimonas oryzihabitans* and *Bacillus cereus*. In From the three *Acinetobacter* sp. strains isolated, dripping waters, gram-negative bacteria producing one grew very fast and produced a large amount of crystals amounted to 15.2% of the population de- crystals surrounding the colonies, which were visible

positive isolates were able to use glucose, fructose, tected, higher than in rock (from 9.8 to 0%). Among maltose and esculin, however, gram-negative isolates gram-positive bacteria, the most common genus

originated by the action of bacteria. The bacteria the ability to produce crystals and therefore could 3.4. *Crystal formation* play a role in the deposition of calcium carbonate polymorphs on the rock surface. Interestingly, Went In the ceiling and walls of Altamire cave some (1969) found a fungus regularly associated with the rmations of calcite, aragonite, and hydromagnesite active tip of stalactites where crystallization of

after 24 h. After one week the crystals were distrib- **4. Concluding remarks** uted all over the plate (Fig. 4). This isolate was only able to use acetate and pyruvate but not citrate or In general, cultural bacterial communities from glucose, the other two strains also used citrate. ground waters have low proportions of gram-positive

face of the most active *Acinetobacter* sp. after 20 from Altamira cave. These communities are mainly days culture for structural and morphological analy- composed of gram-negative rods and cocci (*En*sis. Calcium carbonate polymorphs were identified *terobacteriaceae* and *Vibrionaceae*) while those of by FT-IR spectroscopy (Falini et al., 1996), SEM, rocks were mainly *Streptomyces* spp. (Groth et al., and confirmed by X-ray diffraction. The main crys- 1999). This difference cannot be ascribed to a tals were vaterite (85%) and calcite (15%). Calcite difference in culture media composition (Petrifilm was precipitated by different genera of bacteria (Ben for water and TSA for rock isolation) as water Omar et al., 1997), aragonite by *Acinetobacter* samples were also inoculated directly in TSA with strains (Del Moral et al., 1987). Vaterite is highly similar results. In addition, most of the bacteria unstable and is rarely found. Lowenstam (1981) only isolated have the ability to precipitate crystals in reported vaterite in *Rhodophyta*, in addition to a few vitro and probably in natural habitats. These findings animal taxa. Support the assert by Canaveras et al. (1999) that $\frac{1}{2}$

Acinetobacter sp. strain in B-4 medium. Dark crystals are due to trality, and cyclic nutrient limitations. In addition,

Crystals were collected from the monolayer sur- bacteria. This was also the trend in dripping waters aragonite and hydromagnesite formations in Altamira cave is a biologically-mediated process.

> Actinomycetes are well known for their ability to grow on very poor media (Lechevalier and Lechevalier, 1967) and streptomycetes exist for extend periods as resting arthrospores that germinate in the occasional presence of exogenous nutrients (Goodfellow and Williams, 1983). Dissolved organic carbon content in dripping waters is highly variable, from less than 5 mg C l⁻¹ in winter to about 2200 mg C l⁻¹ in late spring. Dissolved organic matter from soil, which is the origin of the organic carbon found in the dripping waters, contain aliphatic organic acids and phenolic compounds (Saiz-Jimenez and Hermosin, 1999). Guggenberger and Zech (1994) reported that water-soluble organic matter from soils is composed of polymeric lignocellulose degradation products. Both lignocellulose and humic materials are almost selectively degraded by actinomycetes (Crawford et al., 1983; McCarthy, 1987; Ball et al., 1989; Ball et al., 1990; Kontchou and Blondeau, 1992; Dari et al., 1995) and even humic acid is used in an actinomycete isolation medium (Hayakawa and Nonomura, 1987).

Differences in community composition may be attributable to environmental conditions in the cave: high humidity ($> 95\%$), relatively low and stable Fig. 4. (a) and (b). Crystals of calcium carbonate produced by an temperature (around 13°C), water pH close to neuthe addition of a tetrazolium indicator dye. Bar is 0.125 mm. factors influencing attachment of bacteria (microroughness, substratum chemistry and pH, fluid Special Publication 6, Society for Industrial Microbiology, duramic etc.) have to be concidered. All this seems Arlington. dynamic, etc.) have to be considered. All this seems
to favor long-term colonization and selective growth
dependence or calcite polymorphism by mollusk shell macroof actinomycetes on cave surfaces over other hetero-

frophic bacteria present in dripping waters.

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