



## Biological data extraction from imagery – How far can we go? A case study from the Mid-Atlantic Ridge

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### ABSTRACT

In the past few decades, hydrothermal vent research has progressed immensely, resulting in higher-quality samples and long-term studies. With time, scientists are becoming more aware of the impacts of sampling on the faunal communities and are looking for less invasive ways to investigate the vent ecosystems. In this perspective, imagery analysis plays a very important role. With this study, we test which factors can be quantitatively and accurately assessed based on imagery, through comparison with faunal sampling. Twelve instrumented chains were deployed on the Atlantic Eiffel Tower hydrothermal edifice and the corresponding study sites were subsequently sampled. Discrete, quantitative samples were compared to the imagery recorded during the experiment. An observer-effect was tested, by comparing imagery data gathered by different scientists. Most factors based on image analyses concerning *Bathymodiolus azoricus* mussels were shown to be valid representations of the corresponding samples. Additional ecological assets, based exclusively on imagery, were included.

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### 1. Introduction

For the past few decades, deep-sea environments such as hydrothermal vents have been intensively studied by researchers worldwide. These extreme ecosystems are traditionally characterized by their remoteness (both from shore as in depth), and hostile environment (e.g. elevated temperatures, high hydrogen sulphide concentrations, steep chemical gradients). The chemosynthetic micro-organisms present at hydrothermal vents support very specific faunal assemblages. In order to increase our knowledge about the functioning and dynamics of vent ecosystems and their associated communities, faunal sampling is considered fundamental. Information collected through sampling on species composition, densities and biomass is, as such, essential to understand community ecology and biological productivity (Juniper et al., 1998). In this perspective, collecting animals is also crucial to study organism's physiology. With

time, new techniques have been developed, allowing more precise analyses, higher quality samples and long-term studies.

Behind this progress, however, the question of the potential effects of unregulated sampling on these isolated ecosystems arises (Tyler et al., 2005). At hydrothermal vents, sampling is carried out with manned submersibles and Remotely Operated Vehicles (ROV's) which use their robust manipulator arms to sample the uneven and mostly hard substrata inhabited by vent organisms. The irregularity of these sampling surfaces makes quantitative sampling complicated and can result in local perturbations, lastingly changing fluid flow patterns and faunal communities. Such fluid flow modifications were shown to have a profound influence on local assemblages (Hessler et al., 1985, 1988; Fustec et al., 1987; Tunnickliffe, 1991; Sarrazin et al., 1997, 2002; Shank et al., 1998). Despite the precautions taken in choosing more easily accessible sites and relatively flat surfaces, sampling at vents remains intricate and can still be disturbing for faunal communities. This threat was recognized by vent researchers early on (Tyler et al., 2005) but only a few studies have been carried out on sampling impacts. Tunnickliffe (1990) is one of those few studies that assessed sampling effects on deep-sea hydrothermal vents and its fauna and demonstrated that the local vent community, in this case siboglinid polychaetes, had major difficulties to maintain itself after such an

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anthropogenic disturbance, resulting in a lack of subsequent colonization of the sampling area. Overall, while certain geological structures appear to recover quickly from sampling effects, the faunal community, depending on the scale of sampling, may take up to several years to regain its original state (i.e. prior to sampling).

All submersibles and ROV's are equipped with piloting cameras that record imagery footage while diving and carrying out experiments. Contrastingly to sampling, image analysis is a non-invasive technique, which is also one of its greatest advantages. Moreover, imagery analysis permits investigating a larger surface (larger spatial coverage) than discrete sampling and therefore gives a more extended overview of the habitat and faunal communities. It is also more random than sampling, especially in the type of irregular terrain encountered at vents. Imagery analysis has been used to estimate surface areas, faunal coverage and the presence of associated fauna in a variety of ecosystems, such as coral reefs and coastal habitats (Norris et al., 1997; Magorrian and Service, 1998; Ninio et al., 2000). At hydrothermal vents, imagery analysis already proved its value as it is an indispensable, and often the only, tool available in analysing community distribution and temporal variations (Desbruyères, 1998; Sarrazin et al., 1997; Shank et al., 1998; Tunnicliffe et al., 1997; Desbruyères et al., 2001; Tsurumi and Tunnicliffe, 2001; Shank et al., 2003; Copley et al., 2007a,b; Nees et al., 2008; Marcus et al., 2009; Cuvelier et al., 2009, 2011a,b; Podowski et al., 2009; Fabri et al., 2011). Nevertheless, not all parameters are easy to assess based on imagery, and even while video studies are not invasive, they often require ground-truthing with collection of discrete samples at some stage in the investigation (Godet et al., 2010). However, once the connection between what we see and what we sample is established, visual recognition can be used to assess faunal distributions and physico-chemical environmental changes over time and on larger scales (Cuvelier et al., 2011a,b). With all that said, the question remains: how far can we go with imagery analyses without compromising on accurateness?

Currently, hydrothermal vents are somewhat "protected" in the sense that an international code of conduct was drawn up by the international scientific community (Tyler et al., 2005; Devey et al., 2007). The world's first deep-sea Marine Protected Area (MPA) was established in 2003 on the Endeavour segment on the Juan de Fuca Ridge (Canada, Devey et al., 2007). In the Atlantic, the Lucky Strike vent field was proposed as an important conservation area (Azores, Santos et al., 2003) and was included in the Oslo and Paris Conventions network of MPA's in 2007. It was also accepted in 2009 by the European Community as a Site of Community Importance under the Natura 2000 network (O.J. L 344, 2009). The goal of these MPA's is to preserve the integrity of hydrothermal ecosystems including the species and habitats to allow a better understanding of their natural diversity, productivity and dynamics. These protective measures do not allow sampling at certain sites and therefore emphasize the increasing importance of imagery and the accurateness of its interpretations in monitoring natural dynamics.

This paper concurrently investigates the potential and limitations of imagery analyses. In addition, we highlight complementary features that can only be extracted from imagery and are thus an asset to discrete sampling. As part of an in-depth ecological study, twelve instrumented chains equipped with temperature probes were deployed on visibly different faunal assemblages on the Eiffel Tower edifice (Lucky Strike vent field, Mid-Atlantic Ridge – MAR). These twelve sites were thoroughly investigated, comparing imagery analyses with the corresponding ground-truth samples. In order to evaluate possible subjectivity in analysing imagery, we also compared the imagery analyses carried out by two different scientists to evaluate the observer's effect. The major objective of this study is to compare the results obtained by video imagery with those obtained through quantitative sampling.

## 2. Material & methods

### 2.1. Study site

Data gathering for this study was carried out during the MoMARETO cruise (2006) which took place in the MAR region, situated south-west of the Azores Triple Junction (ATJ). Data were collected on the hydrothermal Eiffel Tower edifice, a sulphide structure located south-east of the central lava lake of the Lucky Strike vent field. This 11 m high edifice, situated at a depth of 1690 m, is colonized by *Bathymodiolus azoricus* mussels as well as by *Mirocaris fortunata* shrimp assemblages (Comtet and Desbruyères, 1998; Desbruyères et al., 2000, 2001; Cuvelier et al., 2009). After several monitoring/screening dives, 12 locations were chosen at various places on and around Eiffel Tower (Fig. 1), featuring various assemblages and possibly different physico-chemical conditions. A temporary "chain", equipped with autonomous temperature probes, was placed on each sampling location (Fig. 2).

### 2.2. Faunal sampling

The ROV Victor6000 was used to sample the fauna at each sampling site. First the temporary chain was removed, after which the mobile fauna on the area previously covered by the chain was sampled with the ROV suction sampler. Subsequently, the underlying faunal assemblages (on the same area as previously sampled with the suction sampler) were sampled with Victor's arm grab and put into separate sampling boxes. Finally, a second suction sample was taken on the bare surface in order to recover the remaining fauna. Surfaces sampled have been delineated on the faunal assemblages and sampling locations as presented in Fig. 2.

### 2.3. Image analysis

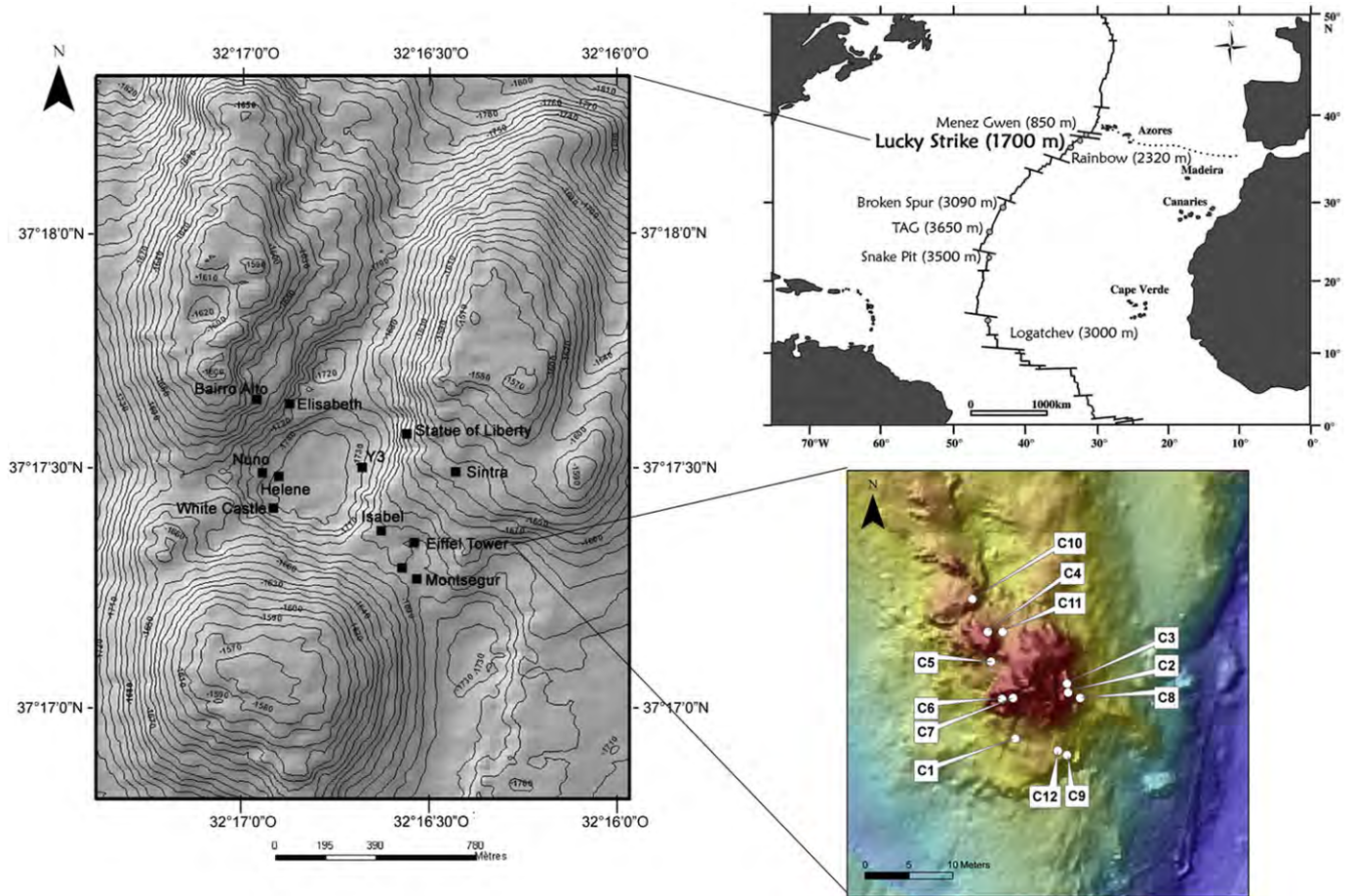
#### 2.3.1. Imagery collection and characteristics

During all the dives, video imagery data was recorded by a 3-CCD camera (HYTEC, VSPN 3000) and a digital high-definition still camera (Sony, Cybershot), mounted above the principal camera of the ROV. Pan, tilt and zoom were kept constant to the extent possible (i.e. not compromising manoeuvres from the ROV). All imagery data was digitally recorded on DVDs. Lighting was provided by 8 flood lights on a fixed bar at the front of the ROV, totalling 5 kW.

For the image analysis, all imagery available (from the approach of the sampling site and the sampling itself) was used, comprising high-resolution photographs, video imagery and screen-stills. Still-images were used as templates to map the surface sampled and analyse the fauna within. Preference was given to the use of high-resolution images, which were mostly available for all sites, however when these were unavailable or unusable, screen-stills were used as a template for analyses. High-definition images had a resolution of 2048 × 1536 pixels while screen-stills were 696 × 576 pixels. Additional high-resolution photographs, featuring zoom-ins, different angles, alongside video imagery from different angles were used to study these sampling sites to reduce the visual distortion of the irregular hydrothermal surfaces, the differences in lighting and shadows cast by the ROV.

#### 2.3.2. Evaluation of sampled surfaces

In order to allow comparisons between the different sampling sites, the surfaces sampled were measured with pixel-based image analysis software IPLAB Spectrum<sup>®</sup> as described in Sarrazin et al. (1997) (Fig. 2). For this study, the twelve instrumented chains with links of 9 cm were used to set the calibration. Because the chains were removed before sampling the fauna, the length of a remarkable



**Fig. 1.** Localization of the Lucky Strike vent field along the Mid-Atlantic Ridge with an overview of the vent field and its bathymetry. In detail, a map of the Eiffel Tower edifice is shown with the locations of the 12 “temporary” instrumented chains deployed during the MoMARETO cruise in 2006 on the Mid-Atlantic Ridge.

feature (mussel, rock) present in the digital photos taken before and after sampling was measured. This feature was then used to calibrate the post-sampling photo and the sampled surface was outlined manually. This was done for each sampling unit separately. In order to reduce the variability due to manual tracing, a mean of three measurements was used to evaluate the final surface area (see Sarrazin et al., 1997 for details). To allow comparison between imagery and the faunal ground-truth samples, we restricted ourselves to analyse what was within these sampled surfaces.

### 2.3.3. Biological data extraction and ground-truthing

**2.3.3.1. Faunal composition and taxonomic richness.** High-definition photographs as well as video footage were analysed to evaluate faunal abundances on the different sampling sites. Taxa were identified to the lowest taxonomic level possible, based on the imagery available. In order to allow comparison in faunal composition between imagery and corresponding faunal samples, organisms from the actual samples were assembled in similar higher taxonomic groups. Abundances were subject to parametric or non-parametric pair-wise testing between imagery and sampling, depending on the normality of the data. Besides this, taxonomic richness was compared between sampling and imagery. Both sample and site-based expected taxonomic richness were assessed by using rarefaction curves (Gotelli and Colwell, 2001; Gauthier et al., 2010).

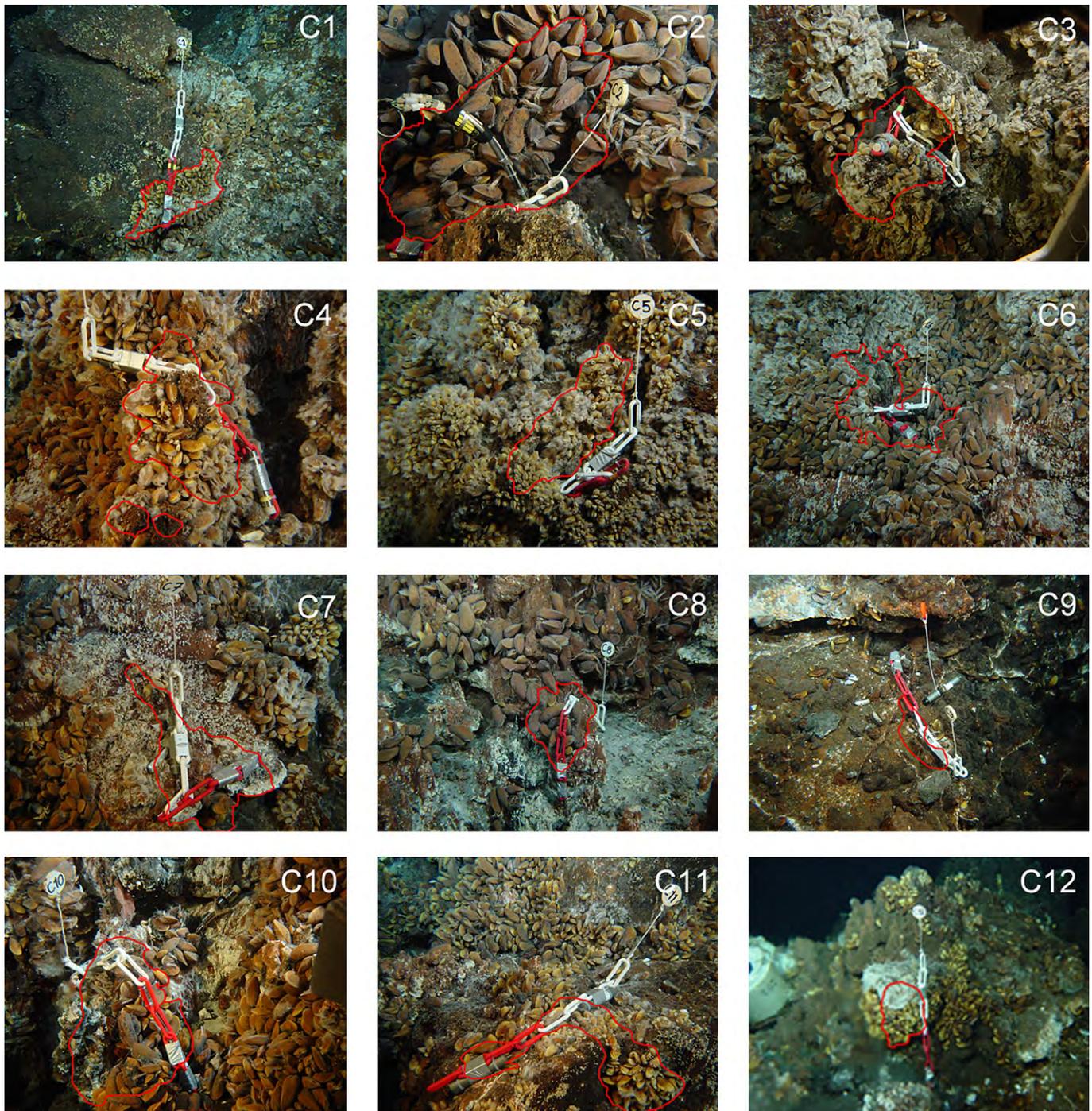
**2.3.3.2. Mussel lengths and biomass.** Mussel sizes were analysed on-screen. The entire length of the animal was measured when

possible but most often, only the width was measurable. In those cases, the lengths were estimated based on a strong relationship between widths and lengths ( $Y = 0.8438x - 0.0241$ ,  $R^2 = 0.9519$ ,  $n = 621$ ), as extracted from the ground-truth samples (Sarrazin et al., unpublished results). Using a paired  $t$ -test, mean mussel sizes as measured on-screen were compared with those from the corresponding ground-truth samples. The evaluation of the lengths also enabled the determination of biomass of mussels on imagery, based on the strong relationship linking shell length to biomass measures, also deduced from the corresponding ground-truth samples (Sarrazin et al., unpublished results). Based on this dataset, Ash Free Dry Weight (AFDW,  $R^2 = 0.9889$ ) and Wet Weight without shell (WWws,  $R^2 = 0.9892$ ) were calculated and used in the biomass estimations carried out here.

### 2.3.4. Additional ecological factors based exclusively on imagery

**2.3.4.1. Mussel valve opening.** The aperture of the shells (or valve gape) was also analysed as the number of mussels that had their valve opened, in order to assess the amount of individuals possibly displaying filtration activity, inhalation of sulphide-rich vent fluids, or gas exchange with the environment.

**2.3.4.2. Microbial coverage.** The microbial cover was measured directly on-screen, by tracing the different surfaces covered by the white filamentous mats visible on the image. The surface coverage of the microbial mats was calculated as a mean of three measurements to assess the final microbial surface area (see Sarrazin et al., 1997 for the image analysis method).



**Fig. 2.** Overview of the 12 instrumented chains (sampling units) and the delineation of the surface sampled, showing the assemblages and the habitats. Each link of the chain is 9 cm long. C12 is a bit blurry due to rising shimmering water in front of the camera lens.

### 2.3.5. Observer's effect

For the first six sampling units, an observer effect was evaluated. The imagery data were analysed by two scientists and differences between their observations were assessed. Observations subject to observer effects' tests were (i) surfaces sampled, (ii) mussel densities, (iii) mean mussel size, (iv) percentage of mussels with microbial cover and with opened valves and (v) abundance of associated fauna. Mean mussel size for C3 was only estimated by one observer, as mussels at this site were covered by microbial mats and single individuals were hard to discern. Differences were analysed with paired *t*-tests when values were normally

distributed. If the normality assumption was still not met after transformation, a Wilcoxon test for paired samples, which is the non-parametric equivalent of the paired samples *t*-test, was carried out.

## 3. Results

### 3.1. Comparing data extraction from imagery with samples

All biological data extracted from the imagery (video and photo) are listed in Table 1. Faunal densities and abundances were assessed

**Table 1**  
Overview of the biological data extraction based on imagery compared to that from sampling for corresponding taxa. The estimated size of the surface sampled as well as the densities of the visible fauna, mussel size and biomass, valve openings and microbial mat covering are given. \* contains macrofaunal taxa that were not detectable on imagery and are thus not listed in the table. The mean temperature (with standard deviation) on each sampling unit was measured and was added for information purposes solely.

Chains	C1	C2	C3	C4	C5	C6	C7	C8	C9	C10	C11	C12
Surface sampled (m <sup>2</sup> )	0.062 ± 0.004	0.070 ± 0.012	0.082 ± 0.00	0.049 ± 0.001	0.049 ± 0.004	0.077 ± 0.007	0.01 ± 0.004	0.028 ± 0.002	0.013 ± 0.003	0.061 ± 0.006	0.033 ± 0.002	0.032 ± 0.002
<b>Densities (ind/m<sup>2</sup>)</b>												
<b>From Imagery</b>												
<i>Bathymodiolus azoricus</i>	2226	1029	1098	1918	4592	883	700	536	0	1246	1727	3938
Alvinocarididae	177	529	207	449	41	338	7900	3536	462	1164	697	656
<i>Segonzacia mesatlantica</i>	32	29	24	20	0	104	700	107	0	16	152	31
Polychaeta	48	0	0	20	20	0	0	0	0	0	30	31
Ophiuridae	16	0	0	0	0	0	0	0	0	0	61	125
Gastropoda	387	0	0	224	204	0	400	0	923	0	909	0
<b>From sampling units</b>												
<i>Bathymodiolus azoricus</i>	5484	857	2049	3776	7857	792	1200	214	7077	1475	4939	8500
Alvinocarididae	323	1743	573	449	0	1338	10,900	3500	0	1426	697	125
<i>Segonzacia mesatlantica</i>	0	71	12	0	0	39	0	107	0	0	152	0
Polychaeta	5661	1000	4341	8592	9041	403	500	250	0	1197	6242	13469
Ophiuridae	0	0	0	0	0	0	0	0	0	0	30	63
Gastropoda	2516	43	610	796	1143	26	1300	143	3154	0	4485	1031
<b>Macrofaunal taxon richness</b>												
From Imagery	6	3	3	5	4	3	4	3	2	3	6	5
From sampling units*	10	7	13	7	9	7	6	7	4	5	14	12
<b>Mussels and microbial cover</b>												
Mean Mussel size (mm)	36.99 ± 8.91	70.96 ± 14.22	47.84 ± 13.85	42.36 ± 15.51	29.36 ± 11.36	57.32 ± 9.47	46.29 ± 5.25	73.1 ± 15.62	N/A	60.28 ± 14.30	45.21 ± 12.72	42.08 ± 12.54
Mussel AFDW (kg/m <sup>2</sup> )	0.54	0.89	0.42	1.07	0.48	0.49	0.31	0.59	N/A	1.74	0.58	1.06
Mussel WWws (kg/m <sup>2</sup> )	3.67	7.09	3.04	7.54	3.12	3.73	2.21	4.73	N/A	13.34	4.18	7.43
Valve openings %	24.64	19.44	4.44	85.11	67.11	32.35	14.29	40.00	N/A	11.84	42.11	3.17
Microbial surface cover (m <sup>2</sup> )	0.009 ± 0.00	0.00 ± 0.00	0.032 ± 0.002	0.012 ± 0.002	0.013 ± 0.001	0.009 ± 0.00	0.00 ± 0.00	0.00 ± 0.00	0.00 ± 0.00	0.00 ± 0.00	0.001 ± 0.00	0.012 ± 0.00
Microbial surface cover %	15.0	0.0	38.7	23.4	27.3	11.3	0.0	0.0	0.0	0.0	3.8	39.5
<b>Temperature (°C)</b>	4.91 ± 0.47	6.50 ± 1.66	5.35 ± 0.33	5.67 ± 0.50	5.11 ± 0.48	6.04 ± 0.59	5.39 ± 0.30	7.49 ± 1.54	4.79 ± 0.12	8.79 ± 2.71	4.85 ± 0.26	4.80 ± 0.33

based on high-definition photographs and video footage from the twelve chain deployment sites. Species were grouped into higher level taxa, in order to avoid identification errors based on the imagery. Some species were identifiable to species level, e.g. *B. azoricus* and *Segonzacia mesatlantica*. For the others, we chose to group them into family levels or classes (Table 1). An overview of the ground-truth sample data (Sarrazin et al., unpublished results), corresponding in taxonomic level to those obtained with the video imagery, is also given (Table 1).

### 3.1.1. Faunal composition and diversity

*B. azoricus* mussels were the main constituent of the assemblages sampled, followed by *M. fortunata* shrimp. In order to allow ground-truthing, the same grouping in taxa was applied to the data originating from the ground-truth samples (Table 1). Generally, for polychaetes and gastropods and, to a lesser extent for *B. azoricus* and alvinocaridid shrimps, a higher abundance was discernable in the ground-truth samples when compared to imagery (Fig. 3). This observation was very pronounced for the polychaetes and the gastropods (Fig. 3). Exceptions for mussels were C2, C6 and C8 and for shrimp C5, C8, C9 and C12 that featured higher densities on the imagery. There were no *B. azoricus* visible directly on the sampled surface of sampling unit C9 but they were observed in the surroundings (Fig. 2). These faunal abundances were tested pairwise (Wilcoxon test) between sampling and imagery for each taxon and showed significant differences for mussels ( $V = 72$ ,  $p$ -value = 0.007), polychaetes ( $V = 0$ ,  $p$ -value = 0.004) and gastropods ( $V = 0$ ,  $p$ -value = 0.004) but no significant differences for shrimp ( $V = 12$ ,  $p$ -value = 0.13). Ophiuridae and *Segonzacia* crabs, on the other hand, showed higher abundances and were encountered more often on the imagery than in the samples, be it in a non-significant way ( $V = 30$ ,  $p$ -value = 0.10 for the crabs and  $V = 6$ ,  $p$ -value = 0.17 for the ophiuroids).

Overall, when investigating taxonomic richness, several macrofaunal taxa present in the ground-truth samples were not found on the imagery. These include small-sized species belonging to polychaetes and gastropod taxa, but also pycnogonids, ostracods, actinids, halacarids, nematods, copepods, tanaids and amphipods. For all locations, the taxonomic richness is higher in the samples, than that recognized on the imagery. The maximum taxonomic richness for the imagery analyses was found at sampling units C1 and C11, where 6 different taxa were encountered. In the ground-truth samples, the highest taxonomic richness was found in C11 with a total of 14 macrofaunal taxa, followed by C3 with 13 taxa (Table 1). The lowest richness from the imagery was attributed to sampling unit C9 where only 2 taxa were observed on the imagery. In the samples, the lowest taxonomic richness was observed in C9 as well with 4 taxa (Table 1). The taxonomic richness as deduced from imagery compared to that from sampling was significantly different (paired  $t$ -test,  $t = 5.9699$ ,  $df = 11$ ,  $p$ -value = 0.00009). Furthermore, an additional test (ANOVA) confirmed that the results obtained from sampling (S) and Imagery (I) are significantly different from each other i.e. that variation in taxonomic richness between S and I is larger than between the sampling locations ( $F = 20.28$ ,  $df = 1$ ,  $p = 0.0002$ ). However, there was a significant positive correlation between the taxonomic richness's from imagery and sampling ( $t = 2.4654$ ,  $df = 10$ ,  $p$ -value < 0.05), thus implying that they show similar tendencies.

The sample-based rarefaction curves (Fig. 4) also showed that the expected taxon richness for discrete sampling was higher than that for imagery, clearly exhibiting the limits of imagery for estimating richness. Moreover, the curve of the imagery analyses seemed to reach an asymptote rather quickly while that of the sampling still did not yet stabilize for the number of samples investigated here, reinforcing the fact that video imagery is a rather

weak estimator of overall species or taxon richness in these hydrothermal assemblages (Fig. 4).

### 3.1.2. Mussel size and biomass

*B. azoricus* mussels were visibly the most abundant species in the sampling units and were therefore subject to several additional analyses, such as size measurements and biomass estimations (Table 1). There was a difference in mean mussel size between sampling and imagery (paired  $t$ -test,  $t = -1.9922$ ,  $df = 10$ ,  $p$ -value = 0.074) (Fig. 5). The biomass calculations were based on the existing significant relationship between size and biomass for the sampled individuals. No significant differences, not for Ash free Dry Weight (AFDW,  $t = -0.6634$ ,  $df = 10$ ,  $p$ -value = 0.52), nor for Wet Weight without shell (WWws  $t = -0.8894$ ,  $df = 10$ ,  $p$ -value = 0.3947) were observed between the sampled individuals and those estimated from the mussels measured on the video imagery.

## 3.2. Complementary imagery analysis data

### 3.2.1. Valve openings

The proportion of open mussels was also assessed as a percentage of all mussels present in the sampled area. These values varied between 3% and 85% on C2 and C4 respectively (Table 1).

### 3.2.2. Microbial cover

Microbial cover was evaluated as the sampled surface covered by the microbial mats. It varied from 0.001 m<sup>2</sup> to 0.032 m<sup>2</sup> and no microbial mats were observed on sampling units C2, C7, C8, C9 and C10 (Table 1).

## 3.3. Observer's effect

For the first six sampling units (C1–C6), two observers assessed the same parameters, independently from one another. Even though differences between the observers are noticeable (Fig. 6), no significant differences were revealed between the sampled surfaces ( $t = -1.2828$ ,  $df = 5$ ,  $p$ -value = 0.26), mussel densities ( $t = 0.6672$ ,  $df = 5$ ,  $p$ -value = 0.53) and the percentage of mussels with their valve opened ( $t = 0.9695$ ,  $df = 5$ ,  $p$ -value = 0.38). The densities of the associated fauna were systematically higher for Observer 1 than for Observer 2 but these differences were not significant ( $p$ -value > 0.1, Fig. 6).

On the other hand, a significant difference for the mean mussel size estimations was revealed between the two observers ( $t = -3.4232$ ,  $df = 4$ ,  $p$ -value = 0.027), which were systematically slightly higher for Observer 2 (Fig. 6). The mean mussel size of C3 was not taken into account in the statistical tests because it was only measured by one of the observers. The results also show that, with the exception of C1, the two observers encountered the same number of taxa on all sampling units (Fig. 6).

## 4. Discussion

### 4.1. Biological data extraction

#### 4.1.1. Faunal composition and diversity

When comparing imagery and sampling, some distinct differences in observations tend to come out. For instance, crabs and ophiuroids are more abundant in the imagery data than in the samples, although non-significant. Such a trend could not be revealed for the shrimp. Nevertheless, we can conclude that assessing mobile faunal presence and/or abundance of some larger macrofaunal taxa is more accurate based on imagery than sampling, since mobile organisms tend to escape during sampling. On the other hand, the abundance of smaller organisms such as

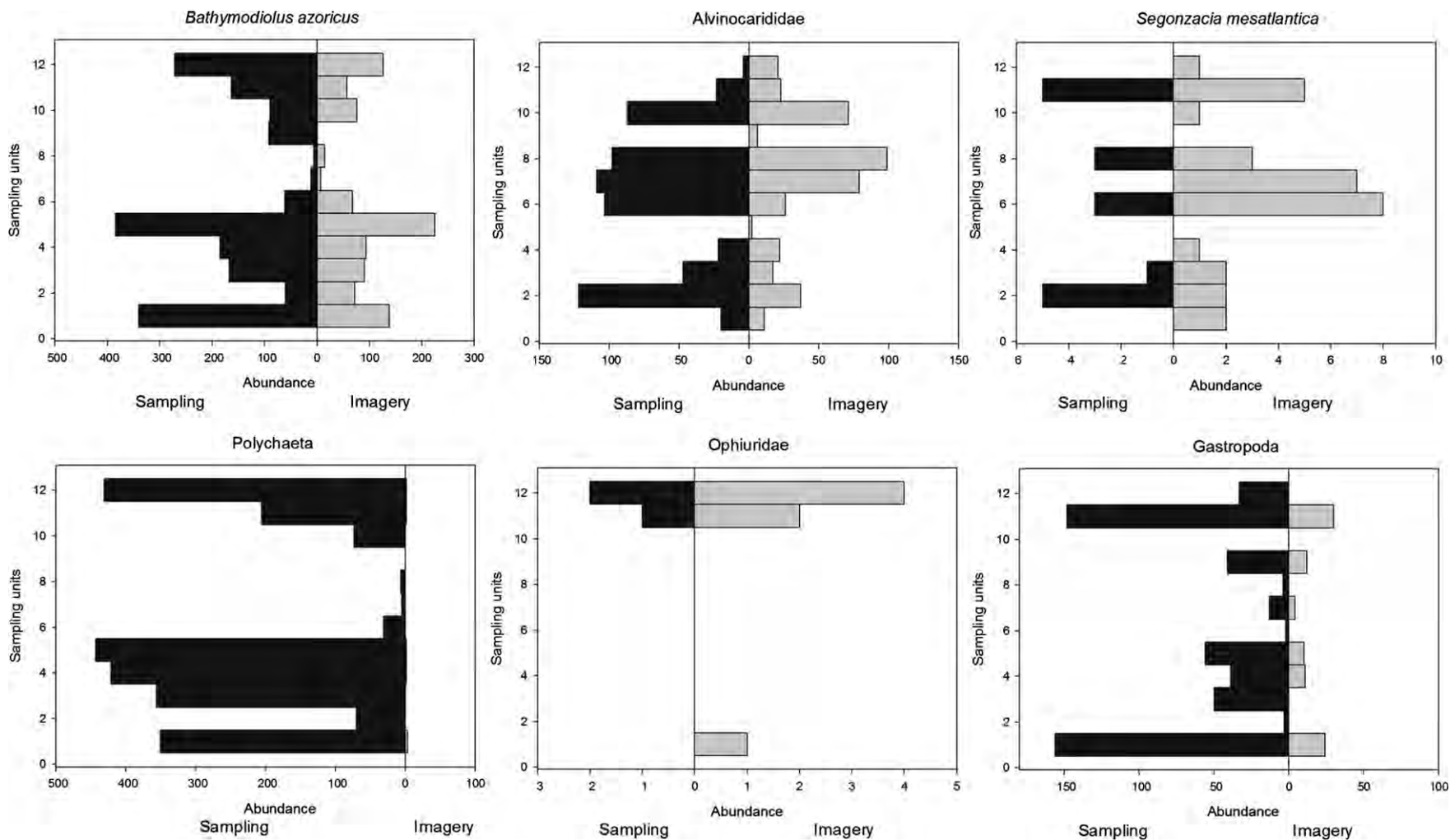


Fig. 3. Faunal abundances as identified on discrete sampling (left = black) and imagery (right = grey).

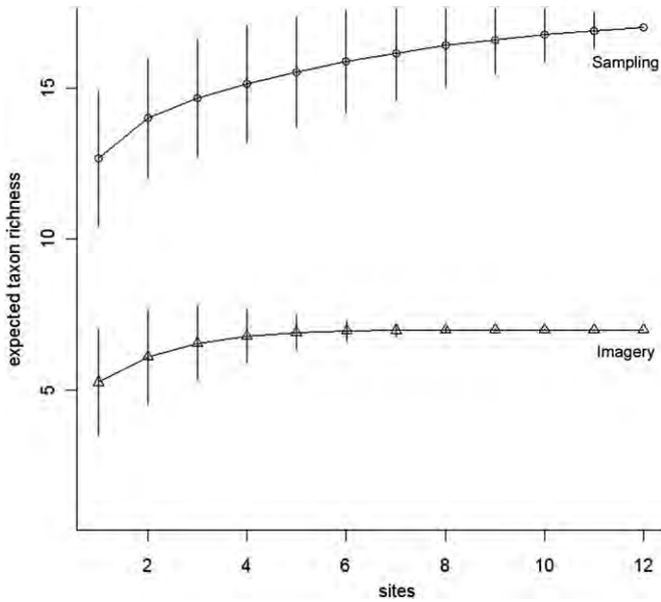


Fig. 4. Rarefaction curves displaying the expected taxonomic richness in the sampling units and imagery.

polychaetes, amphipods, copepods and gastropods, which tend to live in interstitial spaces between mussels (Van Dover and Trask, 2000), or even inside the mussels' mantle cavity (e.g. the polynoid polychaete *Branchiopolynoe seepensis*, Britayev et al., 2007), is almost impossible to evaluate on imagery and therefore explains the significant differences in abundances and taxonomic richness between imagery and samples. The same goes for younger and therefore smaller organisms. For example, there were no mussels visible in the image analyses from sampling unit C9 while 24 individuals were found in the samples (Sarrazin et al., unpublished results). These were not seen on imagery, most likely due to their small sizes (<1 cm).

Taking all of the above into account, the diversity measures based on imagery represent a clear underestimation of the real diversity occurring on the Eiffel Tower hydrothermal edifice. At best, 71% of the real taxonomic diversity was assessed with imagery data while at worst, the estimation goes down to 23%. This is confirmed by the sample-based expected taxonomic richness curves, in which the curve for imagery reaches an asymptote at a smaller number of samples than the curve related to sampling, which does not level off. Finally, as larger groups of taxa are considered, the taxonomic richness assessed is most likely to only represent the minimum species richness of the Lucky Strike vent community (Gauthier et al., 2010). However, the positive

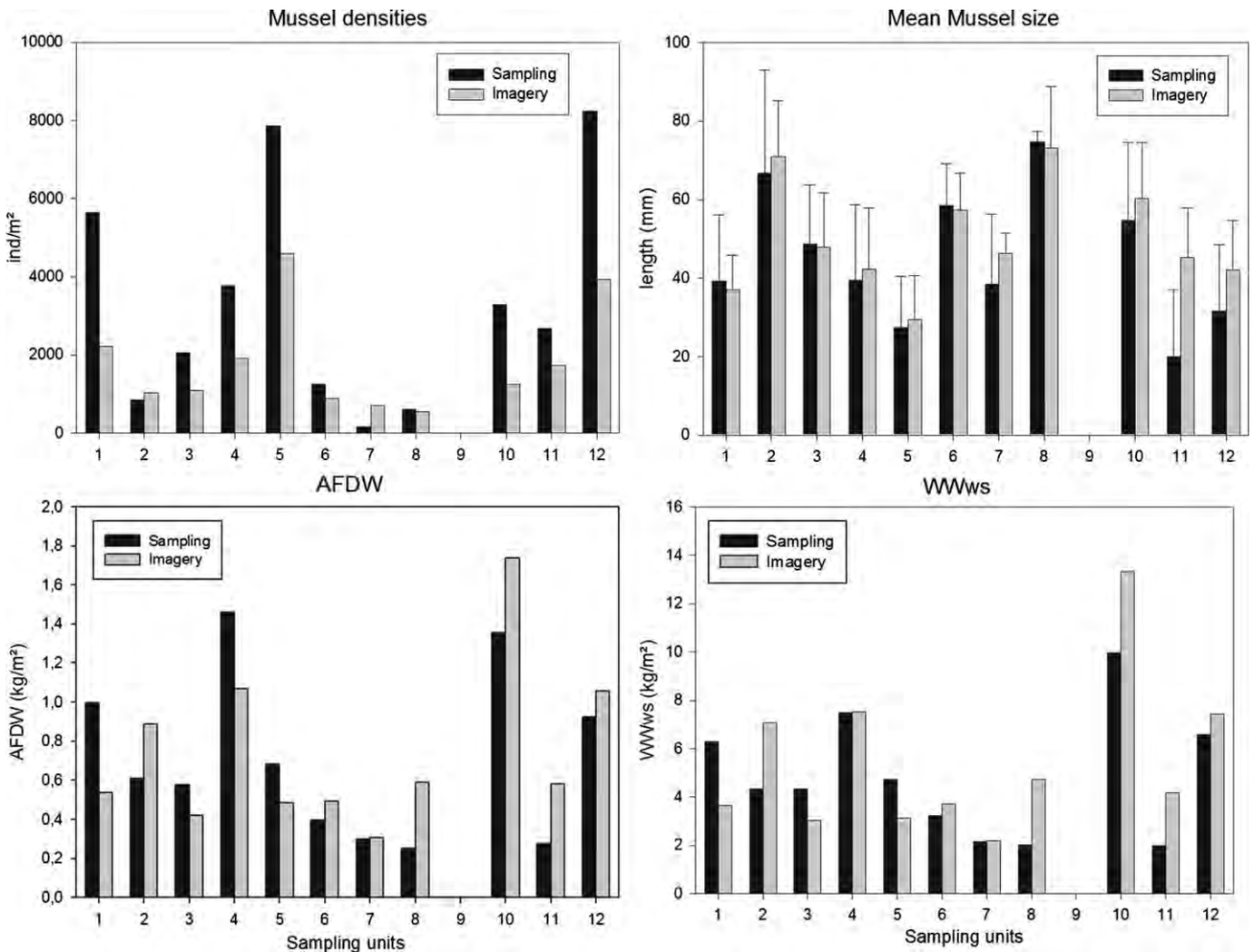


Fig. 5. Comparison of *Bathymodiolus azoricus* mussel densities, mean sizes and biomass (wet weight – WWws – and ash free dry weight – AFDW) between the samples and video imagery.

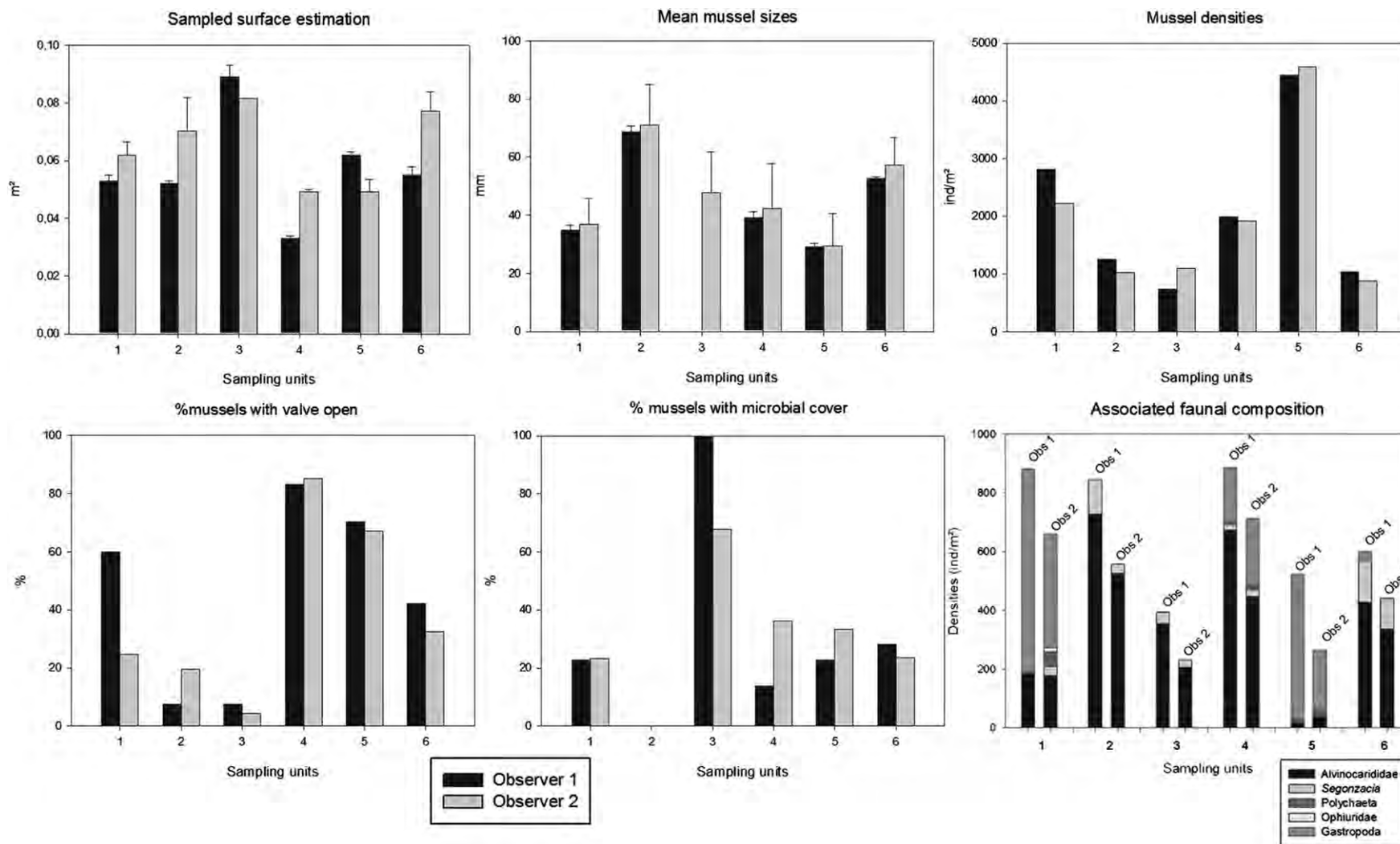


Fig. 6. Illustration of the “observer effect” on different estimations: sampled surface, mussel size and density, percentage of opened mussels and composition of associated fauna.

correlation in taxonomic richness between sampling and imagery shows that researchers can reveal trends in taxonomic richness or diversity occurring between the sampling sites based on imagery alone.

#### 4.1.2. Densities, size and biomass

Densities (number of individuals/m<sup>2</sup>) are extrapolated from the sampled surface estimations. This is an indispensable part for imagery analyses, but at the same time it represents the biggest challenge, as this comprises 2D analyses of 3D surfaces. In addition, mussels are known to stack up to several layers (Johnson et al., 1994) and offer secondary surfaces for associated organisms (Van Dover and Trask, 2000). For example, limpets and other gastropods living on mussel shells and bare surfaces can be difficult to distinguish and identify because of their shape and colours (camouflage). This is also the main explanation for the significant differences in abundances/densities between imagery and sampling. On the other hand, the transport of faunal samples from the sampling units to sampling boxes can be tricky and can lead to loss of individuals, which, in several cases, can explain higher densities on imagery as it is the case for mussels on C2, C6, C8.

**Mussel Sizes** – Mean mussel sizes, as measured on imagery, do not show significant differences with the mussels measured in the lab. When investigating size–frequency distributions, it becomes clear that mussels with lengths between 15 and 60 mm are more predominant in the samples than on the imagery, while this size range should normally be visible on the imagery. Evidently, the mussel bed 3D structure is likely causing an underestimation of the visible abundance. Nonetheless, the lengths of the mussels measured on-screen can be considered a valid representation of the reality as shown for fish by Harvey et al. (2003).

**Biomass** – By measuring the mussel lengths on screen, biomass was calculated through the relationship established between the shell length and biomass from the ground-truth samples. No significant differences were revealed between the biomass assessed on imagery and that from the ground-truth samples. This implies that based on the lengths measured on-screen, our biomass estimations appear to be quite accurate, making the mussel length a good proxy for biomass estimations at larger scales. A similar exercise was carried out at Edison seamount (off Papua New Guinea), in which samples featuring vesicomysid clams were analysed by measuring lengths and widths and consequently using this biometric relationship to calculate biomass at a larger scale based on video-transects (Stecher et al., 2003). While hydrothermal vents are known for their high biomasses in the deep-sea (Sarrazin and Juniper, 1999; Van Dover et al., 2001; Govanar et al., 2004; Dreyer et al., 2005), scientists are still struggling with assessing biomass efficiently and correctly in these remote habitats. This is mainly due to the irregular topography, the patchy distribution of vent animals and difficulties in quantitative sampling (Chevaldonné and Jollivet, 1993). Based on the data presented here, the use of biometric relationships can thus be considered a powerful tool to estimate biomass on imagery from remote marine ecosystems. For now, we suggest that at vents it should be systematically validated with discrete samples taken simultaneously to the imagery until it is proven that it can be extrapolated in space and time.

#### 4.1.3. Valve opening

The number of mussels with their valves opened was evaluated. This is only achievable through image analyses, which makes it an extra ecological parameter exclusive to imagery. Valve openings can be an indication of filtration activity in coastal bivalves and is often assessed through measurements of the gape observed between the valves (Riisgard et al., 2003; Maire et al., 2007).

However, in the chemosynthetic vent ecosystem it can also be an indication of endosymbiont exposure to vent fluids or gas exchange. Despite the presence of a double endosymbiotic association with chemosynthetic bacteria in the gills of *B. azoricus* (Duperron et al., 2006), these mytilids are also capable of filter-feeding (Page et al., 1991; Tunnicliffe, 1991; Colaço et al., 2002) which may allow them to survive some time after vent disruption (Fisher, 1995; Copley et al., 1997). It was hypothesized that larger mussels would be more dependent on chemosynthesis, while smaller individuals would depend more on filter-feeding (Martins et al., 2008), but no such relationship could be revealed through our analyses. This could be due to the possibility that for the vent mussels, valve opening behaviour is linked with the general intake of energetic sources be it particles (filtration activity) and/or chemicals (chemosynthesis, respiration). Further research on valve gape activity of vent mussels would be interesting; in particular to evaluate potential links between valve openings and environmental conditions such as hydrothermal activity and hydrodynamic factors. The use of autonomous video camera from deep-sea observatories gives access to such data and could also enlighten us about the potential impacts of submersibles on the evaluation of animal behaviour in the deep-sea.

#### 4.1.4. Microbial cover

The total surface covered by the microbial mats is also only quantifiable through imagery analyses (Cuvelier et al., 2011b) and is thus an asset to ground-truth sampling. Microbial mats are quite abundant in the vent environment but not much is known about their composition, or about the environmental conditions they thrive in. On Eiffel Tower, large areas covered by mussels are mat-free while other *B. azoricus* assemblages are completely covered (Cuvelier et al., 2009). A study by Crépeau et al. (2011) showed a highly diverse microbial community within the microbial mats at Lucky Strike, covering hydrothermal deposits and *B. azoricus* individuals. While the relationships between mussels and mats are still poorly understood, it does not appear to point to a negative one as mussel assemblages covered by microbial mats coexist with mussel assemblages free of microbial mats, both of them being healthy. A commensal relationship thus appears to be the most convincing scenario in which sulphur and methane oxidizers benefit from fluid dispersion by mussels and numerous heterotrophic microorganisms degrade the organic material released by the mussels (Crépeau et al., 2011).

#### 4.1.5. Observer effect?

For most features tested (sampled surface, mussel densities, percentage of mussels with opened valves, densities of the associated fauna), no significant differences were found between the two observers. Only the mean mussel sizes were significantly different. This may be due to overall image quality or to the presence and thickness of microbial coverage (as seen on C3) which may render the basic calibration and/or measurements in general more difficult. A better training of the observers or a clarification of the method used (tutorial) may be considered to reduce such errors/bias. Automated recognition and contouring, using thresholds or pixel-based colour detection, could be helpful to resolve this problem. Additional issues with the image resolution and/or creation of shade casted by the ROV lights on the organisms could lead to different perceptions and measurements on the sampling units analysed.

#### 4.2. Imagery vs. sampling

Both methodologies appear to be indispensable and complementary one to another. Although imagery traditionally needs

**Table 2**

Overview of ecological factors that can be quantitatively assessed based on imagery and sampling at hydrothermal vents. A selection of two recent publications is given as example, with a preference for the MAR regions. This table is an illustration for the differences, advantages and disadvantages of both sampling and imagery, not a review/overview of all literature/examples available. P indicates when imagery was found to be a poor estimator of the proposed variable in the present study. \* are references not from the MAR region.

Imagery	Refs.	Sampling	Refs.
<b>Faunal composition and diversity</b>			
Larger (>1 cm) dominant macrofauna abundances (P)	This study	Including the smaller faunal fraction such as meiofauna, densities for all fractions	Sarrazin et al., 2006*, Zekely et al., 2006; Copley et al., 2007b*
Faunal assemblage distribution Diversity (P)	Cuvelier et al., 2009, Podowski et al., 2009* This study	/	Van Dover and Trask, 2000; Tsurumi 2003*, Sarrazin et al., 2006*
Biomass	Chevaldonné and Jollivet, 1993*; Stecher et al., 2003*, this study	Biomass	Govenar et al., 2004*; Dreyer et al., 2005*
<b>Behaviour and interactions</b>			
Valve opening for mussels	Maire et al., 2007*(non-vent), this study	/	
Retraction-expansion rates for tubeworms	Tunncliffe, 1990*, Chevaldonné and Jollivet, 1993*	/	
Growth	Urcuyo et al., 1998*, Urcuyo et al., 2003*	Length–Frequency distributions to define cohorts/classes	Comtet and Desbruyères, 1998, Thiébaud et al., 2002*
Movements/Locomotion	Bates et al., 2005*; Grélon et al., 2006*	/	
<b>Coverage</b>			
Faunal coverage and surface estimations	Sarrazin et al., 1997*, Cuvelier et al., 2009, 2011b	/	
Microbial mats	Cuvelier et al., 2011b, this study	/	
<b>Environmental variables/hydrothermal activity</b>			
Fluid flow debit	This study, Sarrazin et al., 2009	Fluid flow debit	Ramondenc et al., 2006*; Sarrazin et al., 2009
Local currents	This study	Large-scale currents	Khripounoff et al., 2000, 2008
Fluid relocation and re/de-activation	Copley et al., 1999; Cuvelier et al., 2011b	/	
<b>Temporal variations</b>			
Long-term variations of faunal distribution	Shank et al., 1998*; Copley et al., 2007a; Cuvelier et al., 2011b	Long-term variations in faunal samples	Johnson et al., 2006*, Copley et al., 2007a
Short-term variations of communities (time lapse)	Tunncliffe 1990*; Copley et al., 2009	/	

ground-truthing to corroborate what is seen on-screen, it can be used self-sufficiently afterwards for a number of features, for example to investigate assemblage distribution and variations over time (Cuvelier et al., 2009, 2011a,b). Even to such an extent that, at vents, temporal variation studies are carried out almost solely based on imagery, as it is often the only long-term monitoring tool available. Since an increasing number of deep-sea MPA's will be created and more restrictions on sampling will be imposed, imagery can be a good alternative as it also allows in-depth ecological evaluations, whose value cannot be underestimated. Imagery also completes the data gathered by the samples with information on animal behaviour, fluid flow quantification and coverage of faunal assemblages or microbial mats (Table 2). All these factors contribute to a more profound knowledge and understanding of the vent environment which we cannot have based on sampling alone.

## 5. Conclusions and perspectives

Based on our findings, several biological features deduced from imagery were confirmed to be accurate assessments of ground-truth samples. For instance, the mytilid biomass estimations based on on-screen length measurements were shown to be a valid representation of the sampled biomass and mussel lengths. Moreover, a couple of features appear to be more precise when analysed on imagery than when based on sampling. This is true for the presence and abundance of large mobile fauna such as crabs and ophiuroids that can escape from sampling and also for the microbial coverage which can only be quantitatively evaluated on the imagery.

However, faunal composition, abundance and richness are clearly underestimated when limited to imagery, though taxonomic richness assessed in imagery was shown to reveal similar trends as the ground-truth samples. Nevertheless, sampling is required to correctly evaluate these important biological factors. Besides this, imagery analysis also has multiple assets, with several variables being exclusively available through imagery, such as behaviour, surface and microbial coverage, etc. Moreover, once the structure of the faunal assemblages is well characterized in a region, video imagery can be used to monitor community distribution, dynamics and temporal changes at large spatial scales.

Mean mussel size was the only factor for which the two observers showed significant differences. The development of a clearer protocol or the use of automated calculations could help resolving the observed discrepancies. In the mean time, several scientific teams started developing software featuring automated techniques for imagery analysis for different marine habitats (Aguzzi et al., 2011; Teixido et al., 2011), including deep-sea hydrothermal vents (Aron et al., 2011). On the longer term and with the use of systematic approaches, imagery will represent an essential tool to evaluate the impacts of global warming and anthropogenic activity in the deep ocean. It should be used to help proposing management policies in these remote habitats.

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