# **Chapter 3 Intertidal Death Assemblages as Proxies of Marine Biodiversity. An Example from Northern Patagonia, Argentina**



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**Abstract** Marine conservation biologists have identified mollusks as one of the appropriate surrogate taxa for characterizing marine benthic diversity. In turn, live/dead comparison studies have overwhelmingly demonstrated that mollusk remains are faithful proxies of the mollusk composition of the living communities from which they come, with positive consequences for the paleoecological evaluation of fossil assemblages. In this contribution, we evaluate the way in which mollusk biodiversity is distributed along the lower intertidal to supratidal (high water mark) dead shell assemblages accumulated on a northern Patagonian rocky shore, in order to explore the usefulness of these assemblages as paleontological proxies and potential surrogates of regional biodiversity. A diversity gradient from the lower intertidal to the supratidal was identified which is probably associated with vertical transport, although the influence of gradients of the living community should be tested to confirm this. The outstanding result of this study is the discovery of high levels of diversity among dead shells (31 bivalves and 39 gastropod species) in a single locality and with a moderate sampling effort. The supratidal death assemblage has higher species richness than expected, possibly caused by stranding of the fauna after storms. Nevertheless, this level shows the lowest level of evenness and a strong bias when samples are not sieved through a fine mesh. The record of marine benthic diversity in death assemblages is a promising area of research that deserves to be explored in depth.

**Electronic supplementary material** The online version of this chapter

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#### **3.1 Introduction**

Dead shells accumulated on the sea floor contain a wealth of information which is useful either for assessing important questions of the genesis of fossil deposits (taphonomy) or for studying living communities. In the search for evidence to determine how representative the fossil record is of communities that lived in the geological past, the discipline of taphonomy has developed tools which provide highquality information of living ecosystems. This is achieved by allowing their features to be explored on longer timescales, beyond those typically used by ecologists (Kidwell and Tomašových [2017;](#page-24-0) Olszewski and Kidwell [2007;](#page-24-1) Tomašových and Kidwell [2009a;](#page-24-2) Archuby et al. [2015;](#page-22-0) De Francesco et al. [2013;](#page-23-0) Yanes et al. [2008;](#page-25-0) Hassan et al. [2018\)](#page-23-1). Current developments go still further: it is now possible to identify the effect of human impact on ecosystems, by studying the differences between impacted living communities and time-averaged assemblages accumulated over the past decades or centuries (Erthal et al. [2011;](#page-23-2) Kidwell [2008;](#page-24-3) Yanes [2012;](#page-25-1) Dietl et al. [2015\)](#page-23-3). The relevance of this paleobiological information, which offers us an otherwise inaccessible long-term perspective of biodiversity and community change, has given rise to a new discipline: conservation paleobiology (Barnosky et al. [2017;](#page-23-4) Louys [2012;](#page-24-4) Rick and Lockwood [2013;](#page-24-5) Dietl and Flessa [2011;](#page-23-5) Dietl et al. [2015;](#page-23-3) Kidwell [2009;](#page-24-6) Kidwell and Tomašových [2013\)](#page-24-7).

Biodiversity is of fundamental importance to ecology because it is the consequence of how organisms in communities respond to biotic and abiotic factors (Olszewski and Kidwell [2007\)](#page-24-1). The use of biological surrogates (i.e. estimators, such as polychaetes, crustaceans, mollusks, etc.) to evaluate marine biodiversity is a useful practice in conservation biology because it helps overcome the difficulties inherent in surveying benthic communities: time and cost, the occurrence of undescribed species and the problems of species identification (Tyler and Kowalewski [2017;](#page-25-2) Magierowski and Johnson [2006;](#page-24-8) Mellin et al. [2011;](#page-24-9) Warwick and Light [2002\)](#page-25-3). Research focuses on finding appropriate surrogates for the different types of marine communities and their spatial and temporal variations. Mollusks, which are among the groups selected as appropriate surrogates of marine benthic communities, leave abundant mineralized dead remains, which have been proved to be good proxies of the communities from which they derive (Tyler and Kowalewski [2017;](#page-25-2) Kidwell [2008;](#page-24-3) Smith [2005\)](#page-24-10).

Assessing how diversity transfers from living communities (life assemblages, LAs) to death assemblages (DAs) is a crucial step towards a better interpretation of diversity in fossil assemblages; this knowledge will also help us to evaluate death assemblages as faithful proxies of living communities. The step from LAs to DAs represents the first filter that modifies diversity measurements, through differential

transport and destruction by waves, currents and wind and time-averaging (Archuby et al. [2015;](#page-22-0) Tomašových and Kidwell [2009a,](#page-24-2) [2010a\)](#page-25-4).

In the absence of strong reworking of former beds, such as in the case of ravinement, marine beds encompass a short time span and their skeletal content is considered representative of the average composition of successions of communities along hundreds or, at the most, thousands of years (within-habitat time-averaging of Kidwell and Bosence [1991;](#page-24-11) see also Fürsich and Aberhan [1990\)](#page-23-6). Recently, the quantitative knowledge of the differences between death assemblages and living communities, and the sources of these differences has been greatly improved (Olszewski and Kidwell [2007;](#page-24-1) Tomašových and Kidwell [2009a,](#page-24-2) [b,](#page-25-5) [2010a,](#page-25-4) [b,](#page-25-6) [2011;](#page-25-7) and many more).

# **3.2 Death Assemblages, Taxonomic Diversity, and Taphonomic Fidelity**

Due to the time-averaged nature of DAs, their species composition is not particularly influenced by the short-term species composition fluctuations of living communities (Fürsich and Aberhan [1990;](#page-23-6) Tomašových and Kidwell [2010a;](#page-25-4) Archuby et al. [2015\)](#page-22-0). These short-term fluctuations, such as the local extinction of the surf clam *Mesodesma mactroides* on the Atlantic coasts of Uruguay and Northern Argentina (Fiori and Cazzaniga [1999;](#page-23-7) Dadon [2005\)](#page-23-8), might give totally different results in samples of living communities separated by only a few weeks. However, in this respect, DAs are highly informative due to their inertia in the face of such fluctuations. Compared with living assemblages, DAs which have accumulated over a few decades to several centuries are expected to have an increase in alpha diversity, a decrease in beta diversity (due to spatial mixing), reduced species dominance and increased frequency of rare species (Tomašových and Kidwell [2010a\)](#page-25-4). Additionally, the ecological information of current ecosystems does not span more than a few decades into the past (Rick and Lockwood [2013\)](#page-24-5). If we consider that human occupation of Patagonia dates from around 17,000–14,000 years BP (Perez et al. [2016\)](#page-24-12), baseline ecological studies might fail to identify the non-impacted conditions when assessing anthropogenic influence, since the impacts were already there.

In turn, death assemblages are used to characterise not only the average species compositions of source communities, but also biotic interactions such as local level predator-prey relationships (e.g., Visaggi and Kelley [2007;](#page-25-8) Yanes and Tyler [2009;](#page-25-9) Gordillo and Archuby [2012,](#page-23-9) [2014;](#page-23-10) Martinelli et al. [2013;](#page-24-13) Tyler et al. [2014;](#page-25-10) Archuby and Gordillo [2018\)](#page-22-1), and to compare these along geographical gradients (e.g., Kelley and Hansen [2007;](#page-23-11) Visaggi and Kelley [2015;](#page-25-11) Martinelli et al. [2013\)](#page-24-13). Quantifying predator-prey interactions in living communities implies sampling strategies that are complex and expensive, while the records from death assemblages are a significant source of information.

Studies on taphonomic fidelity (correlation of living and death assemblages) have been developed in marine, freshwater and terrestrial environments (e.g., Fürsich and Flessa [1987;](#page-23-12) Kidwell and Bosence [1991;](#page-24-11) Yanes et al. [2008;](#page-25-0) Tietze and De Francesco [2012;](#page-24-14) Terry [2010;](#page-24-15) more references in Archuby et al. [2015\)](#page-22-0). Studies of marine death assemblages are abundant, although they are mostly based on soft-bottom ecosystems (Olszewski and Kidwell [2007;](#page-24-1) Kidwell [2013\)](#page-24-16), and there are few studies of communities inhabiting rocky bottoms (Zuschin et al. [2000;](#page-25-12) Zuschin and Oliver [2003;](#page-25-13) Zuschin and Stachowitsch [2007\)](#page-25-14). Recently, Archuby et al. [\(2015\)](#page-22-0) assessed the taphonomic fidelity of rocky-bottom communities along 1500 km of the Patagonian Atlantic coast, from death assemblages collected at the high-water mark. These authors found a general agreement between life and death assemblages at the biogeographical province level, working with non-sieved, representative samples (handpicked along transects). Besides the regional agreement, on smaller geographical scales DAs tended to cluster together and separated from LAs. So far, there are no detailed studies on the nature of DAs on rocky shores. A better understanding of the provenance of the diversity differences between life and death assemblages in modern environments is also crucial for correctly interpreting fossil assemblages and ecosystems (Olszewski and Kidwell [2007\)](#page-24-1).

### **3.3 Purpose of This Study**

In this study, we evaluate the way in which species richness and evenness of mollusk death assemblages is distributed along the depth gradient, from the lower intertidal to accumulations at the high-water mark, in Punta Mejillón, Northern Patagonia, Argentina. Our two goals are to improve the understanding of DAs as paleontological proxies and to evaluate their usefulness as surrogates of shallow benthic living communities. Punta Mejillón has little human impact due to its distance from the nearest city (the town of San Antonio Oeste, 105 km away), the difficulty getting there (sand dunes often cover the route), the need for a four-wheel drive vehicle to reach the beach, and also because it is located in a natural protected area (see The Study Area, below). We aim to determine whether the DAs coming from the same habitat but accumulated at different depths include specimens of different species in different proportions (i.e., there is diversity partitioning of DAs along the gradient). We also test the effect of sieving versus non sieving on species richness and evenness. Specifically, we aim to answer the following questions: (i) How does DA species composition vary along the lower intertidal to supratidal gradient? Is there diversity partitioning along the depth gradient in the rocky intertidal belt of northern Patagonia? (ii) Are death assemblages from rocky shores appropriate surrogates of benthic biodiversity in northern Patagonian shallow marine communities? Is there a horizon along the lower intertidal to supratidal belts that collects most of the information on the death assemblages? In other words, where is it best to sample? (iii) What is the effect of sieving on the biodiversity record?

### **3.4 Methodology**

#### *3.4.1 The Study Area*

The study was carried out in Punta Mejillón (PM), located in the Caleta de los Loros natural protected area in Río Negro Province, Argentine Patagonia. The place is difficult to access, which minimizes the impact of tourism and human activities on living communities and death assemblages (Fig. [3.1\)](#page-4-0). Punta Mejillón is on the Atlantic coast  $(41^{\circ}00'37'')$  in the San Matías Gulf. The coastline runs approximately from SE to NW, and the intertidal belt is exposed for more than 300 m during low tides (Fig. [3.2\)](#page-5-0). Biogeographically, PM is in the transition zone between the Argentine and Magellanic Provinces and is characterized by a mixture of species from both biogeographical entities (Balech and Ehrlich [2008\)](#page-22-2). In a recent article, Güller and Zelaya [\(2017\)](#page-23-13) mention a surprisingly high level of mollusk diversity in the San Matías Gulf, which they describe as a hot-spot of diversity.

The northern part of San Matías Gulf, where Punta Mejillón is located, is subject to high levels of physical disturbance, consisting of strong winds, high tidal amplitudes (up to around 9 m) which leave large areas of the intertidal belt exposed, high-energy flows during high tide and low temperatures (sea surface temperatures  $10.1-18.9 \degree C$ ) (Bertness et al. [2006;](#page-23-14) Archuby et al. [2015\)](#page-22-0). Due to the high levels of desiccation stress caused by winds, the region is considered an extremely harsh intertidal rocky ecosystem (Bertness et al. [2006\)](#page-23-14), which results in intertidal communities which are strongly organized by physical stress.

### *3.4.2 Sampling*

Sampling was carried out on 29 November 2013 during low tide, between latitudes S  $41^{\circ}$  00' 32" and  $41^{\circ}$  00' 54". Samples were collected at four levels: 1. accumulation



<span id="page-4-0"></span>



**Fig. 3.2** Pictures of the intertidal belt in Punta Mejillón. **a** Upper intertidal. **b** View from the middle intertidal to the coast. **c** and **d** details of the middle to lower intertidal

<span id="page-5-0"></span>of shells at the high-water mark (supratidal or "Supra"); 2. upper intertidal belt (UI); 3. middle intertidal belt (MI); and 4. lower intertidal belt (LI). At each level, several replicates were extracted from the upper 15 cm using a shovel and were pooled together, until completing 15 L of sediment. The replicates were extracted every 10 m along a transect parallel to the coastline. Since the substrate is mostly hard, samples were taken from depressions filled with sediments in the area surrounding the sampling point. In the absence of a suitable place to extract the replicate, the point was skipped, and the sample was taken at the next point. Samples were sieved in the field with a  $10 \times 10$  mm aperture mesh (coarse) above and a 1 mm  $\times$  1 mm aperture mesh (fine) below so that large shells were captured separately from small shells (Fig. [3.3\)](#page-6-0). The coarse mesh sieve retains shells that are visible and was considered as a proxy "hand-collecting method", that was compared with "whole" samples per level (made by the pooling of coarse and fine samples). The 1 mm sieve was used to explore a suitable sampling strategy for rocky-bottom dominated intertidal DAs from the Patagonian Atlantic coast. Kidwell [\(2002\)](#page-24-17) suggested that sampling with mesh sizes lower than 1 mm might collect a non-representative high amount of larvae and juveniles.

All gastropod and bivalve shells and shell fragments were analyzed and identified to the species level with some exceptions that were unidentifiable due to preservation issues. Other skeletal elements not included in the study were: crab fragments, serpulid tubes, abundant cirriped plates, sea-urchin spines, fragments of bryozoan

<span id="page-6-0"></span>

**Fig. 3.3** Sampling method: sieving samples

colonies, oyster recruits on large valves and polyplacophoran plates. Cirripeds and cirriped plates, although very abundant, were excluded from analysis due to the difficulty in identifying the plates. Gastropod shells and articulated bivalves were assigned one count. Left and right valves of bivalve species were counted separately. The count per species resulted from the sum of articulated specimens plus the most abundant valves (left or right). *Ostrea puelchana* and *Pododesmus rudis* shells that could not be identified were counted together and divided by 2, and then assigned to the respective species. Bivalve fragments were counted if the umbo and at least onethird of the valve were preserved (very small fragments were discarded). Gastropod fragments were counted when they contained the apex and at least half of the shell.

### *3.4.3 Statistical Methods*

Counts were made per level (LI, MI, UI, and Supra). The coarse mesh size fraction of samples was also registered separately for each level. Diversity was estimated using different indices: species richness (S, the raw number of species and by rarefaction), the Shannon-Wiener (H') index, the equitability J index (Hammer and Harper [2006\)](#page-23-15) and the probability of an interspecific encounter (PIE), an evenness index (Hurlbert [1971\)](#page-23-16). Rarefaction to the lowest sample size was calculated in order to evaluate species richness without the effect of sample size. The H' index summarises information on species richness and evenness and correlates with S and sample size, as does the J index. The PIE index was added to obtain an estimation of evenness which was not affected by sample size (Olszewski and Kidwell [2007\)](#page-24-1). Data management

and calculation of the PIE index according to Hurlbert's formula was carried out using standard spreadsheet software. Other diversity indices were calculated using PAST v 3.15 (Hammer et al. [2001\)](#page-23-17).

Samples (levels) were plotted using a non-metric multidimensional scaling (NMDS) ordination analysis to evaluate their similarity. The database was first transformed to percentages per sample, then square root transformed, and then a similarity matrix was calculated based on the Bray-Curtis index (Clarke [1993;](#page-23-18) Clarke and War-wick [2001;](#page-23-19) Clarke et al. [2006\)](#page-23-20). NMDS was carried out using R software, version 3.4.3 (R Core Team [2017\)](#page-24-18).

To test the effect of using samples sieved with coarse mesh (as proxies for collecting by hand), we compared these with the results obtained for whole samples (coarse + fine mesh) by using diversity indices and an ordination plot (NMDS).

Beta diversity was quantified in order to assess both the existence of a gradient along the coastal profile for the four levels (directional turnover) and non-directional variation for comparing the coarse mesh subsample with the whole sample (whole  $=$ coarse plus fine mesh subsamples) (Anderson et al. [2011\)](#page-22-3). To evaluate the gradient in beta diversity, the similarity between the supratidal sample and the samples from all other levels was calculated with the Jaccard similarity index on a presence/absence matrix. The results were plotted in their position on the coastal profile, from Supra to LI. If species turnover along the gradient existed, then a pattern of similarity decrease would be expected from left to right. To determine the differences in species presence/absence in coarse and fine samples, Whittaker's beta diversity index (βw) was calculated between pairs of coarse and whole samples per level, and then plotted in their position on the coastal profile. Higher levels of  $\beta w$  imply a greater mismatch between the coarse mesh samples and the whole samples (Koleff et al. [2003\)](#page-24-19).

#### **3.5 Results**

A total of 12,790 mollusk specimens belonging to 31 bivalve and 39 gastropod species were collected (Table [3.1,](#page-9-0) Supp Appendix A and B). The sample size was uneven between levels due to the variable densities of shells in sediments from the different samples (Fig. [3.4a](#page-8-0)). Lower and middle intertidal samples contained less than half the specimens of the upper and supratidal samples. The coarse fraction per level fluctuated between 17 and 38% (Supra and MI respectively. Figure [3.4b](#page-8-0)). MI, with the smallest sample size (1520 specimens), has the highest percentage of the coarse fraction (38%).



<span id="page-8-0"></span>**Fig. 3.4 a** Size of sample per level. Levels, LI: lower intertidal, MI: middle intertidal, UI: upper intertidal, Supra: supratidal. n: number of specimens. **b** Size of samples per level and proportion of specimens captured in the coarse mesh. **c** coarse mesh. Width of bars express sample-size

# *3.5.1 Alpha and Beta Diversities Across the Intertidal Gradient*

The 70 species identified in this study are distributed differently across samples (Tables [3.1](#page-9-0) and [3.2\)](#page-9-1). The S index is highest in Supra, followed by LI, UI, and MI. However, when standardizing to  $n = 1520$  by rarefaction, the highest diversity is found in LI (46), followed by Supra, MI and UI, which have between 36 and 39 species (Fig. [3.5a](#page-10-0), b). The rarefaction curves show that none of the samples have a stabilizing size pattern (Fig. [3.5c](#page-10-0)), suggesting that larger sample sizes are necessary to accurately document the kind of study.

Evenness differs between levels, and is consistently lowest in Supra, followed by UI, and then LI and MI with higher values (Table [3.2](#page-9-1) and Fig. [3.6a](#page-11-0), b and c). The H' index is highest for LI, while the J index has MI as the evenest sample. The PIE index, which is more reliable for studies with different sample sizes, is highest for LI, followed by MI, UI, and Supra, coinciding with the H' index.

Multivariate ordination using an NMDS plot indicates a similarity between LI and MI, while UI and Supra remain separate (Fig. [3.7a](#page-12-0)). The analysis of beta diversity

<span id="page-9-0"></span>

<b>Tuble of Distribution</b> of counts per fever and mesh size										
	LI		MI		UI		Supra		Total	
Sample type	. C	F	C	F	C	F		F	C	F
n	2031		1520		4095		5144		12,790	
	510	1521	581	939	773	3322	885	4259	2750	10,040
S	49		37		45		55		70	
	37	32	29	31	36	33	34	46	53	52

**Table 3.1** Distribution of counts per level and mesh size

*LI* lower intertidal level, *MI* middle intertidal level, *UI* upper intertidal level, *Supra* supratidal level or high-water mark. *n* number of specimens, *s* number of species (species richness). Sample type: *C* coarse mesh, *F* fine mesh



*n* sample size, *S* species richness, *S*-*rarefaction* species richness at a sample size of 1520, *H'* Shannon-Wiener index, *J* Pielou's evenness index, *PIE* probability of interspecific encounter diversity index

<span id="page-9-1"></span>**Table 3.2** Diversity indices calculated per level and for the pooled sample

<span id="page-10-0"></span>**Fig. 3.5 a** Plot of species richness (S) per level. The bar represents a bootstrap 95% confidence interval. **b** Rarefaction species richness to  $n = 1520$  per sample. The bar includes 2 standard errors. **c** Rarefaction curves per level with 95% bootstrap confidence interval. Species richness on the y axis; sample size on the x axis





<span id="page-11-0"></span>



<span id="page-12-0"></span>**Fig. 3.7 a** Non-metric multidimensional scaling plot between levels. **b** Non-metric multidimensional scaling plot per level and aperture mesh size. **c** coarse mesh sample; W: whole sample (coarse plus fine mesh sample)

allows the identification of a pattern of decrease along the supratidal to the lower intertidal gradient (Fig. [3.9a](#page-15-0)).

# *3.5.2 Effect of Mesh Aperture Size*

The samples sieved with coarse mesh have richness and equitability values which are lower than estimations for whole (coarse  $+$  fine) samples (Table [3.2](#page-9-1) and [3.3\)](#page-13-0). Coarse mesh samples consistently underestimate the species richness of the death



*n* sample size, *S* species richness, *S*-*rarefaction* species richness at a sample size of 511, *H'* Shannon-Wiener index, *J* Pielou's evenness index, *PIE* probability of interspecific encounter diversity index, *C* coarse mesh

assemblage (Fig. [3.8a](#page-14-0)), and are less even for all four levels (PIE index, Fig. [3.8b](#page-14-0)). On the NMDS ordination plot, coarse mesh samples cluster together and are separate from the whole samples (Fig. [3.7b](#page-12-0)). The comparison of Whittaker's beta diversity indices  $(\beta w)$  shows a large mismatch between the coarse mesh and whole samples at the supratidal level (Fig. [3.9b](#page-15-0)), suggesting that at this level the species composition of the coarse sample is the least similar to the whole sample.

#### **3.6 Discussion**

#### *3.6.1 Alpha and Beta Diversity Trends*

There is a general trend in decreasing diversity from the lower intertidal to the supratidal belt, both for species richness and evenness. The pattern is more evident in estimations not dependant on sample size (Figs. [3.5b](#page-10-0) and [3.6c](#page-11-0)) than in measurements associated with sample size (Figs. [3.5a](#page-10-0) and [3.6a](#page-11-0), b). In the case of species richness, its estimation via rarefaction to the lowest sample size  $(n = 1520)$  suggests a trend from LI to UI, although the Supra sample is slightly more diverse than MI and UI (Fig. [3.5b](#page-10-0)). The PIE index shows a decrease in evenness from LI to Supra, and a similar situation can be observed in the J and H' indices, despite the effect of sample size (Fig. [3.8a](#page-14-0), b and c). NMDS ordination does not reflect a clear pattern. However, the values of Bray-Curtis similarity indices between levels follow the LI to Supra gradient (Table [3.4\)](#page-15-1): contiguous samples are more similar to one another than noncontiguous ones. Distance between samples was coded as 1 to 3 (1 for contiguous samples, 2 for LI to UI and Mi to Supra; and 3 for LI to Supra), and the Spearman rank correlation index was calculated between the Bray-Curtis index and distance, thereby obtaining a significant negative correlation of  $-0.93$  ( $p = 0.033$ ). Additional evidence of the influence of the depth gradient in the species composition of samples comes from the evaluation of beta diversity: compared with the Supra level, there is

<span id="page-13-0"></span>**Table 3.3** Diversity indice calculated per level for the coarse aperture mesh sieved samples

<span id="page-14-0"></span>

a decrease in similarity from UI to LI. This is interpreted as a consequence of the depth gradient, whether due to a taphonomic gradient explained by biostratinomic factors (transport by waves and wind, selective destruction), the species composition gradient of the living community, or both. In turn, harsh environmental conditions (high levels of desiccation, strong winds, and wave energy) suggest that the upper intertidal, and particularly the supratidal, belts should have poorer living community diversity; however, this is not seen in the death assemblages, thus inferring vertical transport from the middle and lower intertidal and shallow subtidal. The higher than expected richness in the Supra sample might be a consequence of the trapping (stranding) of shells above the high water mark during energetic storms; shells and live specimens are stranded above the high water mark, and are no longer reached by usual or normal storm waves (López et al. [2008\)](#page-24-20).

<span id="page-15-0"></span>**Fig. 3.9 a** Beta diversity comparison along the supratidal to lower intertidal gradient. D: Jaccard distance index between Supra level and the other three levels. **b** Whittaker beta diversity index between whole sample and coarse mesh sub-sample per level



<span id="page-15-1"></span>



<span id="page-16-0"></span>

Finally, Bray-Curtis similarity was calculated to compare each level with the whole sample (pooled), and it was found that the Supra and UI levels are the most similar to the total sample (Table [3.5\)](#page-16-0). Although in every case abundances were standardized to percentages and square root transformed, the most abundant samples, Supra and UI, might still influence the result and cause this similarity. The best sampling strategy would still be to collect material from every level, but sampling death assemblages accumulated on the high water mark (Supra) level would not lead to important biases. However, it must be considered that the Supra level has the highest bias when fine meshes are not used, as seen below.

Whether diversity along the lower intertidal to supratidal areas of this study follows a gradient of a biological (species composition of the living community), taphonomic (differential transport, destruction and shell production among species) or mixed nature, will hopefully be answered in future studies. As for soft-bottom studies, there is still a need for more actualistic updated research (Olszewski and Kidwell [2007;](#page-24-1) Kidwell [2015;](#page-24-21) Tyler and Kowalewski [2017\)](#page-25-2). Our investigation is particularly relevant because it helps to fill the need for studies of this kind on hard-bottom environments (Smith [2008;](#page-24-22) Archuby et al. [2015\)](#page-22-0).

# *3.6.2 Mesh Size Matters: The Effect of Sieving on Biodiversity Assessment*

There are relevant differences between the coarse fraction and the whole sample (whole samples are composed of coarse and fine mesh samples. For an explanation, see Sampling in Methodology). Coarse samples are less even (Fig. [3.8b](#page-14-0)) and have lower species richness (as expected, since "coarse" samples are part of the "whole" samples of each level; Fig. [3.8a](#page-14-0)). Species richness in coarse samples ranges from 80 to 62% of the values obtained for the whole samples, with equivalent sampling efforts. The differences observed are also reflected in the ordination plot, in which all coarse samples are clustered together and separated from whole samples (NMDS, Fig. [3.7b](#page-12-0)). Discordance in terms of beta diversity is more marked in Supra than in the rest of the samples. This means that in Supra, the mismatch between coarse and whole samples is highest. This is still more relevant if we consider that many minute species, such as *Phlyctiderma semiaspera*, *Turbonilla macaensis* or *Anachis isabellei*, are present in the coarse samples because they were stuck to mytilid byssus,

which would not otherwise have been sampled by hand. Without these queue-jumper species, the biases of coarse samples would have been even larger. Olszewski and Kidwell [\(2007\)](#page-24-1) detected that the evenness of death assemblages is more similar to live assemblages when samples are sieved with a mesh size finer than 2 mm, which was also the case for species richness. A positive bias in evenness and species richness in coarse mesh samples with respect to fine mesh samples (which are in turn more similar to live assemblages), can be caused by the greater durability of large mollusk shells, lower temporal volatility of adult specimens in living communities, or both (Kidwell [2002;](#page-24-17) Olszewski and Kidwell [2007\)](#page-24-1).

The positive effect of sieving with fine meshes in diversity studies based on DAs is supported by evidence from living communities. In a compilation of mollusk diversity (including bivalves, gastropods, polyplacophorans, scaphopods and cephalopods) in the San Matías and San José gulfs, Güller and Zelaya [\(2017\)](#page-23-13) noted that out of the total 196 species described for the whole area, 61 (31.1%) have a maximum size smaller than 10 mm. As a consequence, almost one-third of the species in the assemblage have a lower probability of being collected, and would perhaps be neglected if fine mesh size had not been used. The most abundant gastropod and bivalve species found by these authors, respectively *Parvanachis isabellei* and *Crenella divaricata*, fit this condition.

# *3.6.3 Sampling Issues: Features of Death Assemblages Along the Intertidal Belt*

Dead shell assemblages differ from living communities due to time-averaging and differential transport and destruction (Kidwell [2001\)](#page-24-23). The effect of tidal regime together with the action of waves on the bottom differ in intensity along the intertidal belt. As a consequence of these differences in the intensity of transport, destruction, and sorting of shells, some variation might be expected in shell density, bioclast size, and vertical transport, which would affect the results of sampling. Shell density varies along the intertidal belt, as evidenced in the individuals counted per sample (Fig. [3.4a](#page-8-0) and Table [3.1\)](#page-9-0). The rarefaction curves indicated that none of the samples were large enough to be representative of species diversity, so more sampling is therefore needed, especially from the lower and middle intertidal levels. The proportion of coarse to fine mesh shells varies little; the exception is MI, but the fact that this level has the smallest sample size could explain the difference (Fig. [3.4b](#page-8-0)). Strong vertical transport of shells in DAs is also evident since the species composition of all the samples includes at least some taxa that are characteristic of subtidal to lower intertidal belts (e.g. *Aulacomya atra*, *Venus antiqua*, some *Buccinanops* species, etc.). On the other hand, these rocky Patagonian shores have little diversity of living fauna in the upper intertidal belt, and almost no marine life higher up, in the supratidal fringe (Bertness et al. [2006](#page-23-14) and personal observations).

# *3.6.4 Mollusk Shell Dead Shell Assemblages as Samples of Living Marine Biodiversity*

Two theoretical frameworks have come together in this study: 1. conservation biology has supplied information on the use of biological surrogates of marine benthic communities (Magierowski and Johnson [2006;](#page-24-8) Smith [2005,](#page-24-10) [2008;](#page-24-22) Mellin et al. [2011;](#page-24-9) Tyler and Kowalewski [2017\)](#page-25-2); and 2. taphonomy, and the new, related discipline conservation paleobiology, have contributed with the assessment of the processes that operate between living communities and accumulations of their remains, for those taxa that bear mineralized or highly durable tissues (Kidwell [2001;](#page-24-23) Tomašových and Kidwell [2009a;](#page-24-2) Dietl and Flessa [2011\)](#page-23-5). Put simply: if a particular taxon is an appropriate surrogate for a living community, and the accumulation of its durable remains or death assemblages are good proxies of the living counterpart of the taxon, then the death assemblages are highly valuable tools as rapid and faithful proxies of the living communities.

Different studies coincide on the point that exhaustively sampling living marine diversity is almost impossible, very expensive and particularly time-consuming, mainly due to the difficulties involved in accessing study sites, poor taxonomic knowledge and the high diversity of marine communities (Warwick and Light [2002;](#page-25-3) Magierowski and Johnson [2006;](#page-24-8) Smith [2008;](#page-24-22) Mellin et al. [2011;](#page-24-9) Tyler and Kowalewski [2017\)](#page-25-2). Besides, results show that at least in some cases (depending on habitat type and spatial scale), mollusks are appropriate surrogates of marine communities (e.g., Smith [2008;](#page-24-22) Tyler and Kowalewski [2017\)](#page-25-2). Dead shell assemblages represent time-averaged relics of the communities they come from. Their differences from living assemblages are explained mainly by their time-averaged nature: they are composed of a mixture of successive communities that lived in the same area and are modified by vertical and lateral transport and other biostratinomic agents (Kidwell [2001,](#page-24-23) [2013;](#page-24-16) Archuby et al. [2015\)](#page-22-0). One of the expectations with respect to the features of death assemblages is an increase in species richness and evenness (Tomašových and Kidwell [2010a\)](#page-25-4). Olszewski and Kidwell [\(2007\)](#page-24-1) demonstrated that on average death assemblages surpass living communities in species richness and evenness, although particular examples might have a different pattern (with little frequency). The only study of live/dead comparisons along the Patagonian Atlantic coast detected a systematic increase in both diversity measurements in every single comparison (Archuby et al. [2015\)](#page-22-0), even when sediments were not sieved (just hand collected) and live and dead samples did not coincide in time or extent (living communities were only sampled from the middle intertidal a few years before the collection of death assemblages).

In order to evaluate to what degree our death assemblages can provide relevant information on regional biodiversity, we compared them with data from surveys of benthic communities. Relevant studies on living mollusk diversity in the San Matías and San José gulfs are summarised in Table [3.6,](#page-19-0) including species richness (discriminating between bivalves and gastropods), the nature of the sample (life or death assemblage), the extent of the sampling area, the sampling effort (in terms of



Table 3.6 Living and death assemblages studied in San Matías and San José Gulfs

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from literature, with no indication of additional sample sizes. *np* not provided in the source literature. Acronyms: *PM* Punta Mejillón, *SJG* San José Gulf, *SMG* San Matías Gulf, *PD* Playas Doradas, *PL* Puerto Lobos, *LG* Las Grutas, *SAB* San Antonio Bay. *Biv*, *Gast*, *Moll*bivalves, gastropods and mollusks respectively,

from literature, with no indication of additional sample sizes. np not provided in the source literature. Acronyms: PM Punta Mejillón, SJG San José Gulf, SMG San Matías Gulf, PD Playas Doradas, PL Puerto Lobos, LG Las Grutas, SAB San Antonio Bay. Biv, Gast, Mollbivalves, gastropods and mollusks respectively,

DA death assemblage, LA living assemblage, Extent area comprised in the sampling, K number of samples, SE sampling effort, n number of specimens

number of samples, *SE* sampling effort, *n* number of specimens

<span id="page-19-0"></span>*DA* death assemblage, *LA* living assemblage, *Extent* area comprised in the sampling,

cost and time of the sampling process) and, when available, sample size. Archuby et al. [\(2015\)](#page-22-0) studied a series of live and dead mollusk assemblages, spanning 1500 km from Punta Mejillón to Puerto Deseado in the South of Patagonia. Their case studies from the San Matías Gulf were numbered 2, 4, 5 (LAs), 6 and 7 (DAs). Avaca et al. [\(2008\)](#page-22-4) provided two databases of living communities, sampled in the North and Northwest margins of the San Matías Gulf. Case study 9 corresponds to an extensive study in the San José Gulf and the closest part of the San Matías Gulf (Zaixso et al. [1998\)](#page-25-15). Recently, Güller and Zelaya [\(2017\)](#page-23-13) published a highly-qualified study with information on mollusk species from the San Matías and San José gulfs, which includes results from their own samples and from an exhaustive bibliographic compilation. The authors used 85 sampling points from the intertidal to a depth of 25 m in the subtidal, and also took some (not detailed) samples from deeper bottoms, obtaining a total of 30,481 mollusk specimens, including empty shells and valves. Their database, and most of their study focused on four main areas which together account for 119 species of bivalves (49) and gastropods (70) for both gulfs (when considering only live species found in their samples). When other studies are added, the species count for the whole area reaches 141 species (60 bivalves and 81 gastropods).

Death assemblages offer relatively high levels of diversity compared to life assemblages (Fig. [3.10\)](#page-21-0). Individual (single place) samples of living communities such as 2, 4 and 5 represent low effort sampling but with very little diversity (up to 6 species). Several-point samples of living communities demand high levels of effort (availability of vessels, complex sampling devices, diving). Case studies 3 and 8 are samples of living communities with 122 and 32 sampling points respectively, where only 21 (case study 3) and 23 (case study 8) species were collected. In these two cases, the sampling area was considered of medium size (narrow fringes parallel to the coast). Case study 9 (Zaixso et al. [1998\)](#page-25-15) is a 120 sample point survey in which a species richness of 61 was collected. Case study 12, taken from Güller and Zelaya [\(2017\)](#page-23-13), is based on Zaxso's data plus additional information. Case studies 10, 11 and 13 are based on multi-point sampling of living communities across wide areas of the San Matías Gulf. Case study 14 represents a synthesis of multipoint sampling and all available published information on mollusk diversity in the San Matías and San José gulfs. Güller and Zelaya [\(2017\)](#page-23-13) and Zaixso et al. [\(1998\)](#page-25-15) sieved their samples with less than 2 mm aperture meshes, while Avaca et al. [\(2008\)](#page-22-4) used 40 mm aperture nets. According to Güller and Zelaya [\(2017\)](#page-23-13), sampling without fine meshes drastically reduces diversity, since they detected that more than 45% of species have shells with a maximum size smaller than 15 mm. This difference in sampling strategy might have caused the reduced diversity record in Avaca's samples. Güller and Zelaya [\(2017\)](#page-23-13) consider that the lower than expected diversity found in the San José Gulf compared with their results in other areas is due to the different sampling methods used (they did not actually sample this gulf, but instead summarised information from other studies).

The three death assemblage case studies, 1 (this study), 6 and 7 (Archuby et al. [2015\)](#page-22-0), are single point samples that represent moderate or low effort but offer a relatively high number of specimens and high diversity. Case studies 6 and 7 sampled



<span id="page-21-0"></span>**Fig. 3.10** Plot of calculated species richness in different case studies in the San Matías and San José gulfs. Order of case studies and acronyms of location sampling areas, as in Table [3.6.](#page-19-0) (1: PM (this study). 2: PM (Archuby et al. [2015\)](#page-22-0) (2). 3: SMG (Avaca et al. [2008.](#page-22-4) NOR). 4: LG (Archuby et al. [2015\)](#page-22-0) (2). 5: PD (Archuby et al. [2015\)](#page-22-0) (2). 6: PD (Archuby et al. [2015\)](#page-22-0). (1). 7: PL (Archuby et al. [2015\)](#page-22-0) (1). 8: SMG (Avaca et al. [2008.](#page-22-4) NOE). 9: SJG, SMG. (Zaixo et al. [1998\)](#page-25-15). 10: SAB Güller and Zelaya [2017](#page-23-13) (3). 11: PL Güller and Zelaya [2017](#page-23-13) (3). 12: SJG Güller and Zelaya [2017](#page-23-13) (3). 13: PD Güller and Zelaya [2017](#page-23-13) (3). 14: SMG Güller and Zelaya [2017](#page-23-13) (3). Empty symbols: life assemblages. Filled symbols: death assemblages. Squares: low sampling effort. Triangles: moderate sampling effort. Circles: high sampling effort. Slashed lines join samples from comparable localities

24 and 21 species respectively, which is similar to the 21 and 23 species in Avaca et al. [\(2008\)](#page-22-4) which required 122 and 32 samples for a similar result. However, those samples were taken without sieving. In our study, a single point in Punta Mejillón, sampled using  $1 \times 1$  mm aperture mesh, detected higher species richness than 122 samples throughout the entire San José Gulf.

This study has confirmed that sampling species richness requires sieving with fine mesh sieves for both living communities (Güller and Zelaya [2017\)](#page-23-13) and their associated death assemblages (Kidwell [2002\)](#page-24-17). In turn, death assemblages give excellent results at equivalent levels of sampling effort, if compared with the sampling of life assemblages. This can be explained by the time-averaging and spatial homogenization of successive communities which accumulated to constitute the death assemblage (Tomašových and Kidwell [2010a\)](#page-25-4).

A word of caution is needed here: the case studies of living communities used for comparisons are mainly based on samples taken from subtidal soft bottoms, while our study was carried out in a rocky-bottom intertidal belt. More quantitative research is needed in order to compare life and death assemblages from equivalent habitats and to evaluate the partitioning of biodiversity along depth gradients and in different types of seafloor in the San Matías and San José gulfs.

# **3.7 Conclusions**

One of the outstanding conclusions of this work is that death assemblages accumulated in rocky-bottom coastal environments are highly informative of the regional biodiversity. In this study, we showed that a single point sample contains more species than almost every study of living communities based on dozens of samples for the same region.

The lower intertidal to supratidal (high-water mark) depth gradient contains dead shell accumulations that reflect a gradient in diversity (a decrease in species richness and evenness), as well as a pattern of species turnover. The gradient in the death assemblages is mainly explained by differential transport upwards, and also by a gradient in species turnover in the living community.

Representative samples of death assemblages must be obtained with the use of fine (up to 2 mm aperture) meshes. Supratidal death assemblages, i.e., shells accumulated in the high water mark, are an acceptable proxy of the whole intertidal to subtidal assemblage if a horizon is to be chosen for sampling. The supratidal is more diverse than expected in terms of species richness, probably due to the supply of shells by strong storms, which were then trapped, out of the reach of normal waves or normal storm waves. In turn, supratidal samples show the highest bias in coase-mesh sieved samples (considered as equivalent to hand-collecting). Results from this study suggest that none of the levels is fully representative, and a sample which pools the different levels is recommended. A live/dead comparison is needed to assess this question in more depth. Exploring the live/dead mismatch in rocky-bottom intertidal environments is useful for improving our knowledge of benthic marine life, and for filling a void in the studies of rocky bottoms, as well as for evaluating the novel idea of conservation paleobiology with respect to the detection of diversity-altered, human-impacted ecosystems.

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