

# The role of tissue-specific microbiota in initial establishment success of Pacific oysters

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## Summary

**Microbiota can have positive and negative effects on hosts depending on the environmental conditions. Therefore, it is important to decipher host-microbiota-environment interactions, especially under natural conditions exerting (a)biotic stress. Here, we assess the relative importance of microbiota in different tissues of Pacific oyster for its successful establishment in a new environment. We transplanted oysters from the Southern to the Northern Wadden Sea and controlled for the effects of resident microbiota by administering antibiotics to half of the oysters. We then followed survival and composition of haemolymph, mantle, gill and gut microbiota in local and translocated oysters over 5 days. High mortality was recorded only in non-antibiotic-treated translocated oysters, where high titres of active *Vibrio* sp. in solid tissues indicated systemic infections. Network analyses revealed the highest connectivity and a link to seawater communities in the haemolymph microbiota. Since antibiotics decreased modularity and increased connectivity of the haemolymph-based networks, we propose that community destabilization in non-treated translocated oysters could be attributed**

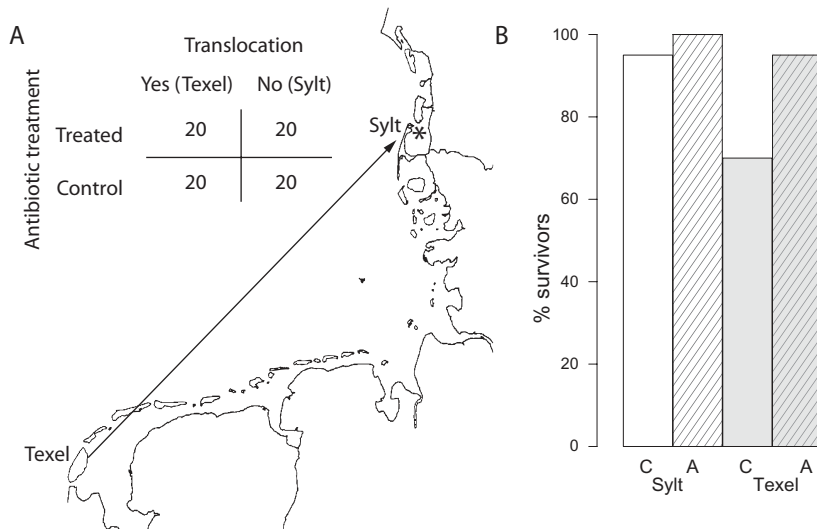
**to interactions between resident and external microbiota, which in turn facilitated passage of vibrios into solid tissues and invoked disease. These interactions of haemolymph microbiota with the external and internal environment may thus represent an important component of oyster fitness.**

## Introduction

Exposure to novel environmental conditions can impose biotic and abiotic stress on organisms (Shea and Chesson, 2002; Hedge and Johnston, 2014). However, such stressors will not only affect the organism directly, but also the microbiota associated with it. Disturbances of fine-tuned interactions among the host, microbiota and the environment may have lethal consequences for the host (Sison-Mangus *et al.*, 2015). Among other functions, resident microbiota are involved in the host immune defence by preventing colonization and establishment of pathogens (Kamada *et al.*, 2013a; Abt and Pamer, 2014; Desriac *et al.*, 2014). While adjustments in community composition can maintain such beneficial functions even under changing environmental conditions, excessive levels of stress can destabilize communities and facilitate shift towards pathogenic states (Rosenberg *et al.*, 2007; Pita *et al.*, 2013; Bauvais *et al.*, 2015; Lokmer and Wegner, 2015). As stability is important for community functioning (Shade *et al.*, 2012), environmental disturbances may bear multilayered consequences beyond the direct effects on the host.

One extreme form of environmental shifts is translocation of organisms to new habitats, within or beyond their normal range. This can happen either unintentionally (e.g. species invasions) or intentionally, like in aquaculture where it is a common practice (Galil *et al.*, 2014). Such habitat shifts will likely be associated with drastic changes in environmental conditions and will not only lead to new interactions of the new environment with the host, but also with its associated microbiota. Further, both the host and its microbiota will not only be affected by changes in abiotic conditions, but will also be exposed to novel biotic environments, including microbes (Jones and Gomulkiewicz, 2012). In humans, for example, travelling to exotic destinations is accompanied by shifts in microbiota composition, potentially leading to health problems (David *et al.*, 2014). While these new colonizers may not harm the

Received 25 May, 2015; accepted 27 November, 2015. \*For correspondence. E-mail alokmer@awi.de; Tel. +49 4651 9564 203; Fax +49 4651 9564 200. Originality significance statement: This is the first study to examine the role of microbiota for the establishment success hosts in a new environment. The host examined is the Pacific oyster, an important invasive and aquaculture species. It also represents the first comprehensive account of Pacific oyster microbiota in the different tissues in order to assess their significance for the oyster fitness and interactions with the environment.



**Fig. 1.** (A) Experimental design showing the number of oysters within each group. The experimental site is marked by asterisk (\*). Experiment duration: 5 days. (B) End-point survival 5 days after translocation and antibiotic treatment ( $n = 80$ ). A, antibiotic-treated; C, control.

organism directly, their interactions with resident microbiota might lead to unforeseeable consequences that cannot be explained by examining the host in isolation (Brown *et al.*, 2009; Koskella *et al.*, 2011; Rillig *et al.*, 2015). To conclusively determine the role of microbiota in colonization success, one needs to investigate the three-way interactions between the host, its microbiota and the environment.

One species that combines several aspects of novel habitat colonization is the Pacific oyster (*Crassostrea gigas*). This species successfully invaded and significantly altered coastal habitats worldwide (Kochmann *et al.*, 2008; Wagner *et al.*, 2012). Moreover, it is a common aquaculture practice to transfer oysters of all life stages between distant cultivation sites (Muehlbauer *et al.*, 2014).

Both transfer and invasion introduce potential mismatches between hosts, their resident, co-adapted microbiota and the novel external biotic environment, including microbes. In other words, the resident microbiota find themselves in a new microbial context, encountering related but locally adapted ecotypes (Martiny *et al.*, 2006; Cohan and Koeppel, 2008) that may alter the microbial community dynamics within the host (Brown *et al.*, 2009; Koskella *et al.*, 2011; Rillig *et al.*, 2015). Despite the applied and fundamental relevance, the consequences of a new microbial context are largely unknown both for the host-associated microbiota as a whole as well as for functionally relevant groups involved.

One important lineage of symbionts and pathogens of marine animals are the bacteria of the genus *Vibrio* (for example, see Hoffmann *et al.*, 2010; Rowley *et al.*, 2014). In oysters, several strains have been implicated in environment-dependent diseases (Garnier *et al.*, 2007; Elston *et al.*, 2008), invoking rapid evolutionary responses

in host populations (Wendling and Wegner, 2015). In addition, the prevalence of pathogenic *Vibrio* spp. in oysters depends on geographic origin (Petton *et al.*, 2015). *Vibrio* spp. and their close relatives of the family *Vibrionaceae* thus represent an ideal focus group to investigate the effects of a new microbial context on the dynamics of opportunistic pathogens.

It is also likely that the disturbance effects will vary among the different tissues of a host. In mammals, for example, gut microbiota are important for immunity and certain changes are related to potentially life-threatening conditions (Kamada *et al.*, 2013b; Ferreira *et al.*, 2014), while the belly-button communities are much less likely to have such an effect (Hulcr *et al.*, 2012). In oysters, haemolymph communities seem to mirror the host condition, as they tend to have low diversity and are dominated by few strains in moribund animals (Garnier *et al.*, 2007; Lokmer and Wegner, 2015). How the microbiota in other tissues relate to oyster condition and to each other is far less known, because most studies have focused on whole body homogenates (Beleneva *et al.*, 2007) or single tissues (Zurel *et al.*, 2011; Trabal *et al.*, 2012; Wegner *et al.*, 2013; Trabal Fernandez *et al.*, 2014). The variation in microbiota composition between the tissues was rarely considered in bivalves (but see Antunes *et al.*, 2010; King *et al.*, 2012), especially in the context of host fitness (but see Meisterhans *et al.*, 2015).

In this study, we set out to experimentally explore how microbiota in different tissues of Pacific oysters are affected by translocation and how the new microbial context affects oyster fitness in a new environment. To do so, we transplanted Pacific oysters from the Southern Wadden Sea (Texel, Netherlands) to the Northern Wadden Sea (Sylt, Germany), and monitored short-term shifts in microbiota composition in different oyster tissues

during the initial phase of establishment in the new habitat (Fig. 1A). The oyster populations on Texel and Sylt have distinct invasion histories and are clearly differentiated by mitochondrial haplotype frequencies (Moehler *et al.*, 2011). In order to distinguish between the oyster- and microbiota-related effects on the establishment success, we treated half of the oysters with antibiotics, thus minimizing the interactions between resident microbes and new colonizers, while the other half was transplanted with their natural resident microbiota. We followed oyster survival and changes in diversity, composition and abundance of oyster-associated bacterial communities as a whole, and *Vibrionaceae* in particular, over the first 5 days in the new habitat. In this way, we could estimate how the composition and diversity of microbiota in different tissues contribute to the successful establishment of Pacific oysters in a new environment.

## Results

### *Oyster survival and immune parameters*

We observed significant differences in survival between the four oyster groups, with only non-antibiotic treated (control) oysters from Texel showing significantly elevated mortality rates (overall test:  $\chi^2 = 12.222$ ,  $df = 3$ ,  $P = 0.007$ ,  $n = 80$ , Fig. 1B, see Table S1 for pairwise comparisons). The mortality could not be linked to genetic differences between the populations (as it differed between the treated and control oysters from Texel, Table S1) nor to differences in the immune system activity, as neither the total haemocyte count (THC) [analysis of variance (ANOVA):  $F_{3,72} = 1.302$ ,  $P = 0.281$ ,  $n = 76$ ] nor the phagocytosis rate per haemocyte (ANOVA:  $F_{3,67} = 0.565$ ,  $P = 0.640$ ,  $n = 71$ ) differed between the four oyster groups. Plasma protein concentration was affected only by antibiotic treatment and thus could not be linked to the mortality either (ANOVA: antibiotic treatment:  $F_{1,71} = 11.314$ ,  $P = 0.001$ , effect size = 0.136; origin;  $F_{1,71} = 0.676$ ,  $P = 0.413$ ; origin  $\times$  treatment:  $F_{1,71} = 0.528$ ,  $P = 0.469$ ,  $n = 75$ ).

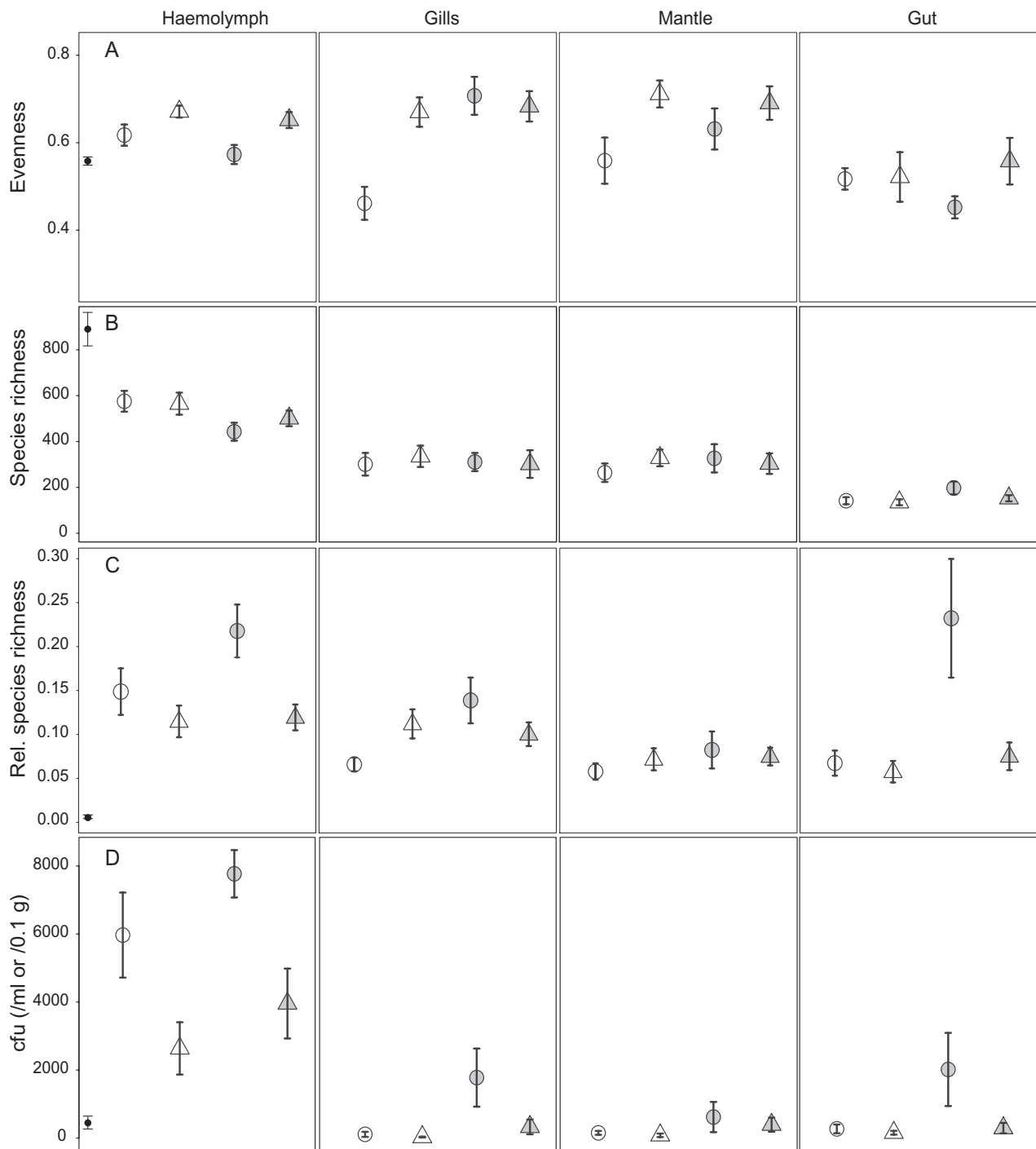
### *Distinction between oyster and seawater microbiota*

Bacterial communities in the seawater differed substantially from those found in oyster tissues. The seawater ( $n = 3$ ) and oyster ( $n = 332$ ) communities were similar in terms of evenness [median: seawater (sw) = 0.549, all oyster samples = 0.641; Wilcoxon rank sum (RS) test:  $P = 0.334$ , Fig. 2A], but overall species richness was much higher in the seawater (median: sw = 873, all oyster samples = 310; Wilcoxon RS test:  $P = 0.008$ , effect size =  $-0.142$ , Fig. 2B). In contrast, the relative operational taxonomic unit (OTU, 97% identity) richness of *Vibrionaceae* was much higher in oyster tissues (median: sw = 0.006, oyster = 0.079; Wilcoxon RS test:  $P = 0.003$ ,

effect size = 0.159), mainly owing to high *Vibrionaceae* diversity in the haemolymph (Fig. 2C). The activity of *Vibrionaceae* – estimated from the colony forming unit (CFU) counts on thiosulfate-citrate-bile salts-sucrose (TCBS) agar – confirmed this pattern, with much higher numbers in the haemolymph than in the seawater (median: haemolymph = 4062.5, sw = 500; Wilcoxon RS test:  $P = 0.004$ , effect size = 0.166, Fig. 2D). All seawater samples were dominated by a handful of taxa belonging to  $\alpha$ -Proteobacteria and *Flavobacteriaceae* (Fig. 3A and B). The most abundant OTUs in the seawater communities (mean relative abundance  $\geq 0.01$ ,  $n = 14$ ) were also found in 85% of the oyster samples albeit in lower abundance, establishing the seawater as a source of bacteria for the oyster microbiome, especially for the haemolymph [mean, median and range of their combined abundance within oyster samples: haemolymph = 0.059, 0.009 (0, 0.433),  $n = 141$ , solid tissues = 0.005, 0.002 (0, 0.083),  $n = 191$ ]. Nevertheless, the seawater and oyster communities were clearly differentiated by community composition (Fig. 3A and B). In addition, non-metric multidimensional scaling (NMDS) plot (Fig. 3A) revealed significantly lower within-group  $\beta$ -diversity of seawater compared to oyster microbiota, which was confirmed by Levene's test for homogeneity of multivariate variances (average distance to median: oyster = 0.614, seawater = 0.201,  $F_{1,333} = 167.9$ ,  $P < 10^{-6}$ , effect size = 0.25).

### *Factors explaining bacterial community differences within oysters: tissue and oyster individual*

Within oysters,  $\alpha$ -diversity significantly differed between the studied tissues, with substantially lower diversity observed in the gut and the highest species richness in the haemolymph (Fig. 2A and B; robust Wilcox bootstrapped ANOVA: evenness:  $F_{3,81.56} = 9.088$ ,  $P < 0.001$ , effect size = 0.368; species richness:  $F_{3,98.493} = 68.052$ ,  $P \leq 0.001$ , effect size = 0.747,  $n = 332$ ). A similar pattern was observed for the relative species richness of *Vibrionaceae* (Fig. 2C; robust Wilcox bootstrapped ANOVA:  $F_{3,127.508} = 7.767$ ,  $P < 0.001$ , effect size = 0.376). Positive correlation between the number of cultivable *Vibrionaceae* and their species richness in the solid tissues [Spearman's  $\rho$ : 0.280,  $P < 0.001$ , confidence interval (CI) = (0.143, 0.427), Fig. S1D] and the lack of such correlation in the haemolymph (Spearman's  $\rho$ : 0.126,  $P = 0.137$ , Fig. S1C) suggest that high *Vibrionaceae* species richness in solid tissues, but not in haemolymph, may indicate a systemic infection. Unlike species richness, the evenness of *Vibrionaceae* was negatively correlated with cultivability in all tissues [Fig. S1A and B; haemolymph: Spearman's  $\rho$ :  $-0.196$ ,  $P = 0.020$ , CI = ( $-0.239$ ,  $-0.021$ ); solid tissues: Spearman's  $\rho$ :  $-0.208$ ,  $P = 0.004$ , CI = ( $-0.347$ ,  $-0.080$ )],

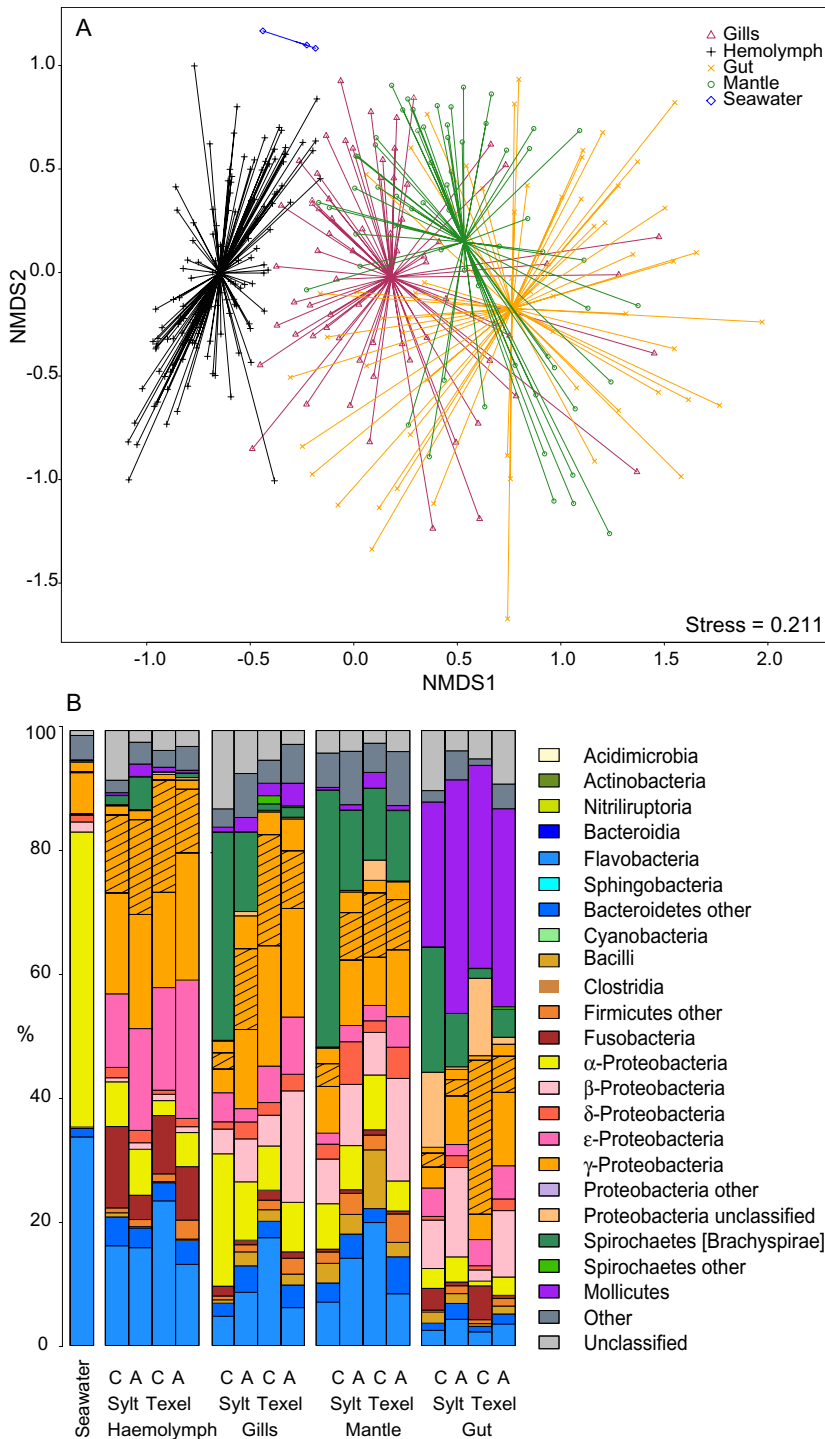


**Fig. 2.**  $\alpha$ -Diversity and *Vibrionaceae* activity in oyster tissues grouped according to oyster origin (clear = Sylt, grey = Texel) and antibiotic treatment (○ = control, △ = treated): (A) overall evenness and (B) overall species richness; (C) proportion of total species assigned to *Vibrionaceae* and (D) cultivable *Vibrionaceae* in 1 ml of haemolymph or approximately 100 mg of solid tissue. Seawater samples (black) are shown in the leftmost plot for reference. Error bars represent standard error of mean.

suggesting that higher overall activity of *Vibrionaceae* tends to be associated with proliferation of few, potentially pathogenic strains.

Tissue also explained a significant portion of variance in community composition, especially when phylogenetic

relatedness was taken into account (compare Bray–Curtis and UniFrac in Fig. 4A), indicating substantial ecological differences between the tissues. Haemolymph communities were clearly distinguished from those in solid tissues by higher relative abundance of  $\epsilon$ -Proteobacteria



**Fig. 3.**  $\beta$ -Diversity and taxonomical composition of seawater and oyster microbiota.

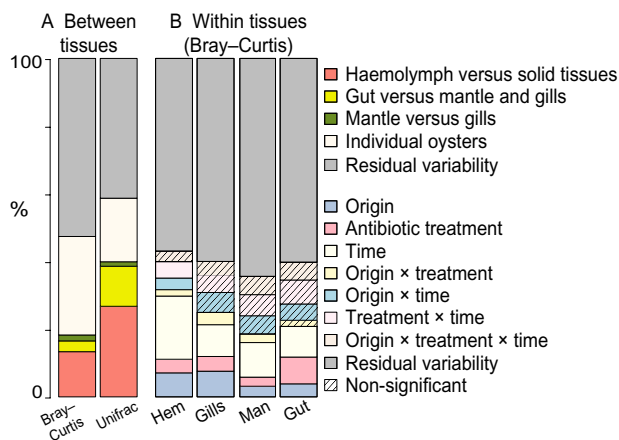
A. NMDS plot of Bray–Curtis distances showing differences between the oyster tissues and the seawater.

B. Mean relative abundance of classes in the seawater and in the tissues, grouped according to oyster origin and treatment. Cross-hatching shows mean relative abundance of *Vibrionaceae*. A, antibiotic-treated; C, control.

(*Arcobacter*), Flavobacteria, Fusobacteria (*Psychrilyobacter*) and  $\gamma$ -Proteobacteria (*Oceanospirillaceae*, *Vibrionaceae*), and fewer Spirochaetes (Brachyspirae) and  $\beta$ -Proteobacteria (Table S2). Among the solid tissues, the most conspicuous difference was the high abundance of Mollicutes (*Mycoplasma*) in the gut, while the mantle and gill microbiota differed only slightly.

In addition to the large effect of tissue, a considerable amount of variability in community composition was explained by differences between individual oysters (Fig. 4A). In this case, however, the explained variability was higher when phylogenetic relatedness was disregarded (compare Bray–Curtis and UniFrac in Fig. 4A).





**Fig. 4.** Variance in community composition explained by (A) tissue type and individual, and (B) oyster origin, antibiotic treatment, time and their interactions in individual tissues. Barplots are graphical representations of Permanova results (cross-hatching shows non-significant factors). Since new oysters were sampled every day, time is partially confounded with between-individual variation.

#### Factors explaining bacterial community differences between oysters: experimental translocation and antibiotic treatment

We examined the effects of population origin and antibiotic treatment on overall  $\alpha$ -diversity and the diversity and activity of *Vibrionaceae* separately for the haemolymph and solid tissues (Fig. 2). The antibiotic treatment increased species richness and evenness of haemolymph microbiota in both oyster populations (Fig. 2A and B, Table 1, haemolymph model, main effect of treatment) and reduced the diversity (Fig. 2C, Table 2, haemolymph model, main effect of treatment) and cultivability of *Vibrionaceae* (Fig. 2D, ANOVA: treatment:  $F_{1,135} = 13.344$ ,  $P < 0.001$ , origin:  $F_{1,135} = 2.509$ ,  $P = 0.116$ , treatment  $\times$  origin:  $F_{1,135} = 0.110$ ,  $P = 0.740$ ,  $n = 141$ ). Interestingly, low evenness of haemolymph microbiota before the transfer to the field correlated with oyster survival (Table 1, haemolymph model, main effect of survival), indicating that disrupted community structure associated with few dominating OTUs may have played a role in the mortalities. In solid tissues, the response to antibiotics depended on the oyster origin. In detail, antibiotics increased the evenness of mantle and gill, but not of gut microbiota in Sylt oysters, while the opposite was true for Texel animals (Fig. 2A, Table 1, solid tissues model, tissue  $\times$  treatment  $\times$  origin interaction). Although neither the treatment nor translocation affected overall species richness (Fig. 2B, Table 1), the antibiotics significantly reduced diversity of *Vibrionaceae* in Texel oysters (Fig. 2C, Table 2, solid tissues model, origin  $\times$  treatment interaction), especially in the gut (Table 2, solid tissues model, tissue  $\times$  treatment interaction). The antibiotics also decreased the number of cultivable *Vibrionaceae*

in Texel oysters, while having little effect on Sylt animals (Fig. 2D, ANOVA: treatment:  $F_{1,181} = 8.227$ ,  $P = 0.005$ , origin:  $F_{1,181} = 12.614$ ,  $P = 0.001$ , origin  $\times$  treatment:  $F_{1,181} = 7.641$ ,  $P = 0.006$ ).

In order to examine the effects of population origin and antibiotics on  $\beta$ -diversity, we analysed each tissue separately (Figs 3 and 4). Despite the high temporal/individual variability, the effects of antibiotics and the population signature remained significant throughout the experiment. Although we observed differences in abundance of some minor genera in the haemolymph (mainly assigned to Flavobacteria,  $\alpha$ - and  $\gamma$ -Proteobacteria), *Vibrionaceae* did not show any significant variation (Table S3). Overall, dominant genera were shared among the oyster groups, but the relative abundance of OTUs belonging to those genera differed. For example, while all haemolymph communities – regardless of treatment or origin – contained bacteria of the genus *Arcobacter*, Texel and Sylt oysters were characterized by a different set of *Arcobacter* OTUs (Table S4).

#### Association networks

The vast majority (89.7%) of associations in the whole-oyster network occurred within tissues (coloured lines in Fig. 5) and only 10.3% occurred between them (grey lines in Fig. 5). Only haemolymph, however, showed higher connectivity than expected by chance (compared with the connectivity of 1000 random node subsets of the same size,  $P < 0.05$ ). We therefore focused on haemolymph for further analysis, and constructed a network for each treatment  $\times$  origin combination to examine microbial association networks in response to antibiotics and translocation (Fig. 6). All resulting networks shared a densely connected area, whose core consisted of the OTUs that were also abundant in the seawater samples ( $\alpha$ -Proteobacteria and Flavobacteria), further establishing the intimate connection between haemolymph and the environment. Other motifs, characterized by *Arcobacter*, *Vibrionaceae* or/and other  $\gamma$ -Proteobacteria and some anaerobes, belonging to Fusobacteria and Clostridia, were recovered in each oyster group as well, and they tended to be negatively associated with the first environmental ('seawater') subcluster (Fig. S2). These shared OTUs had higher than average degree and represented  $23 \pm 5\%$  of nodes within the individual networks (Fig. 6, Table S5). Despite these shared features and although the antibiotic treatment affected only the composition of minor genera within the haemolymph microbiome (Table S3), it significantly altered the association network structure, as antibiotics strongly reduced modularity and increased the connectivity of the networks (Table 3, Table S5). In contrast to control oysters from Sylt, the network based on the control oysters from Texel consisted of more sparsely

**Table 1.** Linear mixed models showing the effects of oyster origin and antibiotic treatment on  $\alpha$ -diversity, namely evenness (clear) and species richness (shaded), in the haemolymph ( $n = 141$ , including before- and after-deployment samples, evenness model fit:  $dAIC = -110.293$ , marginal  $R^2 = 0.211$ , conditional  $R^2 = 0.438$ ; sp. richness model fit:  $dAIC = 34.214$ , mar.  $R^2 = 0.222$ , cond.  $R^2 = 0.308$ ) and solid tissues ( $n = 191$ , even. model fit:  $dAIC = -169.253$ , mar.  $R^2 = 0.310$ , cond.  $R^2 = 0.348$ ; sp. rich. model fit:  $dAIC = 46.934$ , mar.  $R^2 = 0.361$ , cond.  $R^2 = 0.479$ ) separately.

		DF	F <sup>a</sup>	Significant contrasts	Estimate	SE	2.5% CI	97.5% CI
Haemolymph	Main effects							
	Origin	1, 98.237	0.002					
	Treatment	1, 115.947	1.596	Antibiotic versus control	0.032	0.013	0.007	0.057
	Time	1, 84.596	4.704*	Antibiotic versus control	1.210	0.558	0.116	2.303
		5, 91.95	1.277					
	Survival	5, 23.474	1.363	Died versus survived	-0.066	0.023	-0.112	-0.021
		1, 114.134	8.147**					
	Interaction terms							
	Origin × treatment	1, 118.609	1.345					
	Origin × time	1, 66.48	0.012					
Solid tissues	Main effects							
	Oyster	1, 118.131	0.784					
	Bag	5, 92.221	1.012					
	Tissue	5, 116.06	1.134					
	Origin	5, 92.679	0.632					
	Treatment × time	5, 17.966	1.623					
	Interaction terms							
	Origin × time	2, 154.823	11.274***	Mantle and gills versus gut	0.061	0.025	0.025	0.085
	Tissue × origin	2, 110.027	29.572***	Mantle and gills versus gut	0.000	0.000	0.000	1.889
	Origin × treatment	1, 154.983	3.709		0.000	0.000	0.000	0.027
Treatment × time	1, 47.878	0.011		1.817	0.009	0.024	2.479	
Origin × treatment	1, 154.835	13.801***	Antibiotic versus control	0.044	0.012	0.021	0.067	
Time × treatment	4, 52.298	0.424						
Interaction terms								
Tissue × origin	4, 24.652	3.180						
Tissue × treatment	2, 154.658	2.852						
Origin × treatment	2, 109.428	0.411						
Origin × time	2, 154.009	0.310						
Treatment × time	2, 109.434	1.299						
Tissue × time	1, 156.437	2.308						
Tissue × treatment × origin	1, 48.967	3.475						
	4, 154.888	1.038						
	4, 48.415	1.647						
	4, 155.397	2.714*	(Antibiotic versus control): cubic trend	-0.068	0.026	-0.120	-0.016	
	4, 47.871	0.830						
	8, 154.434	1.235						
	8, 108.87	2.150						
	2, 154.265	4.678*	(Mantle and gills: gut): (Sylt versus Texel): (antibiotic versus control)	0.023	0.008	0.007	0.040	
Random variation								
Oyster	2, 110.155	0.166		0.000	0.000	0.000	Inf	
Bag				1.617	0.000	0.000	2.116	
				0.039	0.000	0.000	0.083	
				0.721	0.000	0.000	1.811	

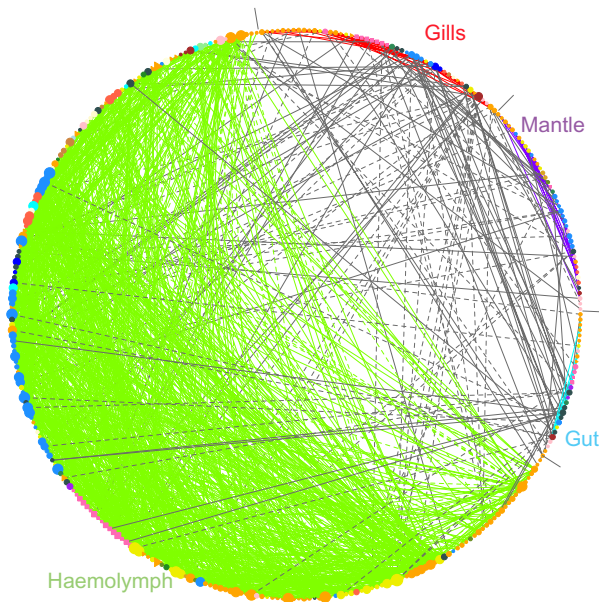
a. Significant levels: \* $\leq 0.05$ , \*\* $< 0.01$ , \*\*\* $< 0.001$ .

**Table 2.** Linear mixed models showing the effects of oyster origin and antibiotic treatment on the relative species richness of Vibrionaceae in the haemolymph ( $n = 141$ , including before- and after-deployment samples, model fit:  $dAIC = -95.295$ ,  $mar. R^2 = 0.211$ ,  $cond. R^2 = 0.577$ ) and solid tissues ( $n = 191$ , model fit:  $dAIC = -206.342$ ,  $mar. R^2 = 0.278$ ,  $cond. R^2 = 0.496$ ) separately.

			DF	F <sup>a</sup>	Significant contrasts	Estimate	SE	2.5% CI	97.5% CI
Haemolymph	Main effects	Origin	1, 82.174	1.190					
		Treatment	1, 86.039	5.951*	Antibiotic versus control	-0.042	0.017	-0.076	-0.008
	Interaction terms	Time	5, 70.55	2.783*	Linear trend	-0.091	0.033	-0.155	-0.027
		Survival	1, 99.267	2.869					
		Origin × treatment	1, 56.625	0.294					
Random variation	Origin × time	5, 74.851	1.988						
	Treatment × time	5, 75.507	0.919						
	Oyster				0.099		0.063	0.133	
Solid tissues	Main effects	Bag	2, 110.837	5.927**		0.035		0.000	0.092
		Tissue	1, 48.861	8.218**	Gills versus mantle	0.029	0.008	0.013	0.046
	Interaction terms	Origin	1, 48.344	1.521	Sylt versus Texel	-0.028	0.010	-0.048	-0.009
		Treatment	4, 18.617	1.138	Antibiotic versus control				
		Time	2, 110.327	2.460					
	Random variation	Tissue × origin	2, 110.475	5.236**	(Mantle and gills versus gut): (antibiotic versus control)	0.016	0.005	0.006	0.026
		Tissue × treatment			(Sylt versus Texel) : (Antibiotic versus control)	0.024	0.010	0.004	0.043
		Origin × treatment	1, 49.798	5.737*					
		Origin × time	4, 48.99	0.985					
		Treatment × time	4, 48.698	0.872					
Random variation	Tissue × time	8, 109.887	2.002						
	Tissue × treatment × origin	2, 111.067	1.574						
	Oyster				0.058		0.026	0.069	
	Bag				0.016		0.000	0.048	

a. Significant levels: \* $\leq 0.05$ , \*\* $< 0.01$ , \*\*\* $< 0.001$ .





**Fig. 5.** Association network showing OTU relationships within and across the tissues. Nodes are coloured by taxonomy (as in Fig. 3) and the size is proportional to the node degree. Full line denotes co-occurrence, the dashed line denotes co-exclusion; edge weight is proportional to correlation strength. Colour of edges connecting OTUs within the tissue correspond to colour of that tissue; inter-tissue edges are grey.

connected modules, which resulted in a pronounced modular structure also reflected by a lower clustering coefficient.

## Discussion

### *Role of microbiota in oyster mortality*

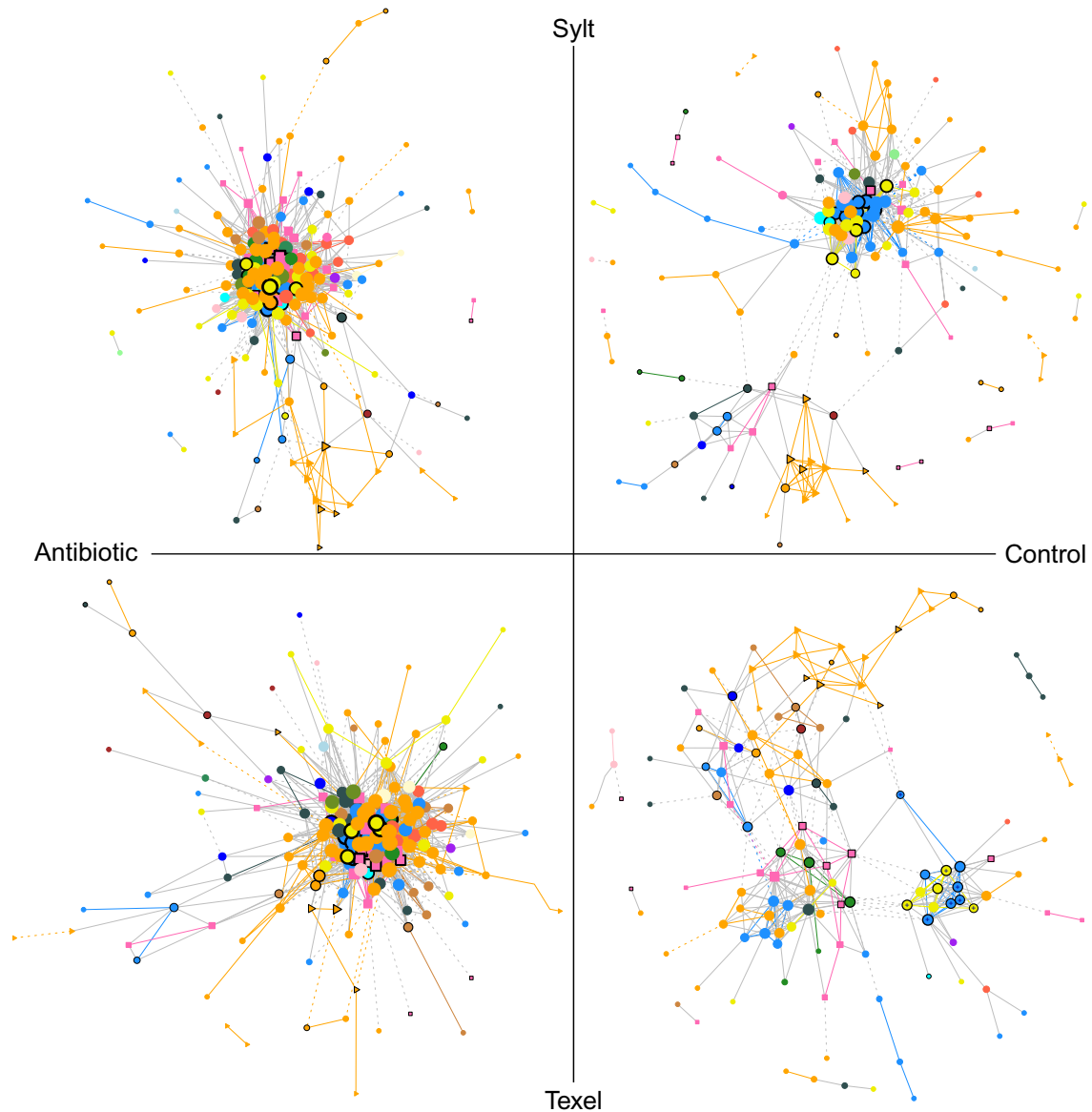
Intentional (transport, cultivation) and unintentional (invasion) translocation exposes plants and animals to novel abiotic and biotic conditions. One example is exposure to new microbes colonizing hosts, where they interact not only with the host but also with its resident microbiota. High mortality of translocated oysters in our study may have been caused by such interactions, since resetting the microbial communities by administering antibiotics prior to the field deployment significantly reduced mortality in translocated hosts. Alternatively, the antibiotic treatment could have mitigated the effects of a transport-stress-induced bacteriosis and affected mortality directly. Since we neither recorded any mortality during transport or in the initial lab-based rearing, nor could we detect any signs of previous disease or significant differences in immune parameters, we consider this to be unlikely. Genetically based direct interactions between hosts and new microbiota could also be excluded as a potential cause of mortality, as genotypes from both oyster populations (Moehler *et al.*, 2011) were randomly distributed

between the antibiotic treatments. The specific change in the network topology – observed in Texel but not in Sylt oysters (Fig. 6) – rather suggests that it was the indirect effects mediated by the resident haemolymph microbiota bacteria (eliminated by antibiotic treatment) and new specific environmental conditions – including the exposure to a new microbial context – that contributed to mortality (Rillig *et al.*, 2015).

Whereas high vibrio load and diversity were observed in the haemolymph of all healthy oysters (including both control groups), they did not translate into high loads in the solid tissues or disease (Petton *et al.*, 2015). High loads of active *Vibrionaceae* in solid tissues were observed only in translocated control animals and could represent signs of systemic infections and thus the proximate cause of mortality. This underlines the importance of performing tissue-specific analyses to understand how the microbiota affects its host.

### *Interactions of haemolymph microbiota with the internal and external environment*

The inter-tissue OTU associations (Fig. 5) and fine-scale individual variation in community composition (compare UniFrac and Bray–Curtis in Fig. 4A) suggest that the oyster microbiome is shaped by genotype (Wegner *et al.*, 2013), condition (Lokmer and Wegner, 2015) and neutral processes (Nemergut *et al.*, 2013). However, the differences in community structure between the haemolymph and solid-tissue microbiota (Figs 2–5) indicate the prominent role of ecological and functional differences between the tissues (Costello *et al.*, 2009; Fuhrman, 2009; Pontarp *et al.*, 2012; Faust *et al.*, 2015a). High network connectivity and the recurring core OTU subclusters (shared groups of OTUs in the networks in Fig. 6 and Fig. S2) imply that a considerable portion of haemolymph microbiota was unaffected by host origin and antibiotics, but rather determined by factors unaccounted for in our experimental design. These subclusters might represent OTU assemblages characteristic for alternative stable community states, possibly related to changes in oxygen concentration associated with periodical valve closing (Sow *et al.*, 2011; Faust *et al.*, 2012). High prevalence of aerobic, seawater-derived bacteria in one subcluster, and the dominance of microaerobic or anaerobic species (Levicán *et al.*, 2014) in the other(s), might reflect a healthy microbiome's response to predictably recurring environmental conditions, i.e. to the tidal cycle (Relman, 2012; Faust *et al.*, 2015b). Shifts in relative OTU abundance within genera (e.g. *Arcobacter*), combined with the persistent community composition at higher taxonomic levels, likely represent yet another mechanism by which the haemolymph microbiota contribute to oyster response to environmental conditions and/or reflect the oyster



**Fig. 6.** Association networks of haemolymph OTUs grouped by oyster origin and treatment. Black frame denotes OTUs shared by all four networks. Nodes are coloured by taxonomy (as in Fig. 3) and the size is proportional to degree. Full line denotes co-occurrence, the dashed line denotes co-exclusion; edge weight is proportional to correlation strength. Edges connecting OTUs within the same class are coloured by taxonomy, else they are grey.  $\Delta$  *Vibriionaceae*;  $\square$  *Arcobacter*. In the last panel, mainly negative correlations between the subcluster with the core consisting mainly of OTUs abundant in the seawater (marked by '+') and the rest of OTUs are apparent (see also Fig. S2).

condition (Lokmer and Wegner, 2015). Overall, the community dynamics of haemolymph microbiota demonstrate its close connection to the internal and external oyster environment – a prerequisite for a role in acclimation and maintenance of homeostasis in highly dynamic environmental conditions.

#### *Haemolymph microbiota in response to translocation and antibiotic treatment*

Time lag between the antibiotic treatment and field deployment resulted in partial recovery and higher diver-

sity of oyster microbiota, likely preventing deleterious effects of depleted microbial communities in the treated animals (Ridley *et al.*, 2012; Abt and Pamer, 2014; Gorokhova *et al.*, 2015). In addition, low evenness of haemolymph communities prior to deployment in the oysters that eventually died suggests that evenness can be used as an indicator for predicting disease (Ransome *et al.*, 2014; Lokmer and Wegner, 2015). Therefore, the diversity of haemolymph microbiota seems to be important for a successful response of oysters to environmental challenges (Ptacnik *et al.*, 2008; Wittebolle *et al.*, 2009; Eisenhauer *et al.*, 2012; Stenuit and Agathos, 2015).

**Table 3.** Properties of the networks depicted in Fig. 6.

	Sylt		Texel	
	Control	Antibiotic	Control	Antibiotic
# Samples	35	36	32	38
# Nodes	149	211	129	190
# Edges	721	2348	362	2742
% Positive edges	82.1	61.1	85.4	56.7
Average degree	9.678	22.256	5.612	28.863
Max degree	49	94	24	93
# Clusters	12	5	7	2
Connectance	0.065	0.106	0.044	0.153
Average path length	3.331	2.693	3.604	2.417
Average betweenness centrality	0.011	0.008	0.015	0.007
Modularity	0.224	0.053	0.599	0.036
Global clustering coefficient	0.625	0.540	0.459	0.565

Detailed comparison with random networks can be found in Table S5.

Apart from diversity, oyster response to the stress exerted by our treatments could have been positively affected by increased co-occurrence network connectivity, which probably reflected higher robustness of haemolymph microbiota against disturbances (Estrada, 2007; Scheffer *et al.*, 2012; Shade *et al.*, 2012). Due to low resolution of 16s rDNA-based analysis (Koeppel and Wu, 2012; Schmidt *et al.*, 2014), we cannot distinguish if the increased connectivity was due to external colonization of the disturbed communities (Robinson *et al.*, 2010) or to a correlated response of resident bacteria to antibiotics. Still, the higher diversity of antibiotic treated communities suggests increased external colonization because antibiotics usually decrease diversity due to dominance of few resistant strains (Manichanh *et al.*, 2010; Dethlefsen and Relman, 2011; Panda *et al.*, 2014), especially when administering a mix of antibiotics like performed here (Pena-Miller *et al.*, 2013).

In control oysters from Texel, high modularity and lower clustering coefficients could indicate a decreased compensation capacity and thus a lower probability to successfully respond to disturbances (Yachi and Loreau, 1999). Although the exact processes behind the observed changes in community structure remain unclear, oysters as filter feeders are in constant touch with a multitude of bacteria, and intense microbial interactions in a new microbial context could have affected community stability (Defer *et al.*, 2013; Rillig *et al.*, 2015). Such destabilization of the haemolymph microbiota, caused, for example, by interactions between closely related but distinct strains (West and Buckling, 2003; Brown *et al.*, 2009; Koskella *et al.*, 2011; Wendling and Wegner, 2015) could explain the spillover of *Vibrionaceae* into solid tissues, resulting in a systemic disease and higher mortality of translocated control oysters.

However, the interpretation of the co-occurrence network topology is not straightforward and the ecological implications vary widely depending on the sort of input

data and the network building criteria (Thebault and Fontaine, 2010; Faust and Raes, 2012; Berry and Widder, 2014; Widder *et al.*, 2014; Williams *et al.*, 2014; Faust *et al.*, 2015a; Peura *et al.*, 2015). In addition, the relationship between the stability and topology of the networks depends on the type of disturbances to which the studied communities are exposed to (Holme, 2011). Therefore, caution must be exerted when linking network structure to the mortalities observed here, unless network characteristics like connectivity and modularity are experimentally manipulated. Eliminating specific bacterial groups by narrow-spectrum antibiotics is one possibility to perform targeted manipulation of the network structure (Rea *et al.*, 2011), and that could elucidate causal links between haemolymph community stability and tissue colonization by pathogens.

#### *Specificity of microbiota in solid tissues*

While high diversity of haemolymph microbiota likely reflects its tight connection to the variable environment, the lower diversity of gut or gill microbiota may reflect the specialized functional roles in target tissues (e.g. such as nutrition in the gut; Duperron *et al.*, 2007; Fraune and Zimmer, 2008; Rodrigues *et al.*, 2010; Fernandez-Piquer *et al.*, 2012). Such tissue-specific functional specialization should lead to increased persistence of microbiota in solid tissues. Here, we mainly observed origin-related persistence of gill microbiota throughout the initial phase of colonization (Table S6). Gills show high bacterial activity in Pacific oysters (Hernandez-Zarate and Olmos-Soto, 2006) and many bivalve bacterial symbionts are situated there (Duperron *et al.*, 2007; Dubilier *et al.*, 2008; Rodrigues *et al.*, 2010). Gill microbiota also exhibits long-term stability (Zurel *et al.*, 2011) and its composition correlates with its host genotype (Wegner *et al.*, 2013). While the differences observed here could represent carryover from the original site (Wegner *et al.*, 2013; Trabal

Fernandez *et al.*, 2014) and would thus gradually disappear (Wendling *et al.*, 2014; Meisterhans *et al.*, 2015), their long-term persistence could reflect differences in the genetic structure and history of both oyster populations (Moehler *et al.*, 2011).

### Conclusion

We experimentally manipulated the interactions of the oyster holobiont with the environment on several levels, including the host genotype (by using genetically differentiated host populations), the resident microbiota (by administering antibiotics) and the external bacterial context (by translocating treated and untreated hosts). Our data highlight the importance to consider microbiota in a tissue-specific context in order to understand the interaction of the holobiont with a newly encountered environment. Specifically, the spillover of *Vibrionaceae* from the haemolymph into solid tissues and a resulting systemic disease seemed to depend on the structure and stability of haemolymph microbiota in response to a new microbial context. This signifies the prominent role of community structure and not necessarily taxonomic composition of the haemolymph microbiome for oyster homeostasis. In order to elucidate the processes behind the observed changes in community structure and to better understand the function of microbiota in different tissues, more studies focusing on the metabolism and physiology of functionally important symbionts within these dynamic bacterial networks are needed.

## Experimental procedures

### Oyster collection and pretreatment

To test interactions between the resident and novel external microbiota, we transplanted oysters from the Southern Wadden Sea to the Northern Wadden Sea and followed their survival and changes in microbial communities over a 5-day period. Southern Wadden Sea oysters ( $n = 40$ ) were collected in de Cocksdorp, Texel, Netherlands ( $53^{\circ} 0' N$ ,  $4^{\circ} 54' E$ ), cleaned of epibionts and transported to the AWI Wadden Sea station on Sylt. The same number of oysters was collected from the transplantation site (Oddewatt, Sylt, Germany,  $55^{\circ} 1' N$ ,  $8^{\circ} 26' E$ ). Initially, the oysters ( $n = 80$ ) were kept in pre-filtered seawater from their original location at ambient temperature ( $\sim 14^{\circ}C$ ). To remove parts of the resident microbiota, we added an antibiotic cocktail ( $100 \mu g l^{-1}$  of each ampicillin, tetracycline, gentamicin and kanamycine, Sigma-Aldrich, Hamburg, Germany) to half of the oysters from each location. The antibiotic concentrations were chosen based on previous tests, where we treated the oysters with the antibiotics and plated out haemolymph on marine agar until no colonies had grown. After 3 days, we took haemolymph samples from the adductor muscle with 23\_1/4 gauge (0.6 dm, 30 mm) needles via notches drilled on the ventral side of the shell immediately after collection (to

give the oysters time to recover). We froze  $\sim 200 \mu l$  aliquots of haemolymph for microbiota analysis at  $-80^{\circ}C$ , and immediately processed the rest of the sample to measure the immune parameters (THC, phagocytosis rate, haemolymph plasma protein concentration). To estimate the number of cultivable *Vibrionaceae*, we plated  $5 \mu l$  of haemolymph on TCBS agar and counted the resulting CFU.

### Experimental set-up and sampling

The pretreatment resulted in four groups, each containing 20 animals: translocated (antibiotic)-treated, translocated control, non-translocated (antibiotic)-treated and non-translocated control oysters (Fig. 1A). For the field transplant, always four oysters (one from each group) were put into a single bag with a mesh size of 1 cm, resulting in 20 bags that were brought out to the original site of collection of the northern Wadden Sea oysters (Odewatt). For the following 5 days, we randomly collected four bags every day. The period of 5 days was chosen because we wanted to focus on the initial phase of the establishment, where we assumed the most drastic and influential changes in the microbiome would occur based on the previous infection experiments (Lokmer and Wegner, 2015). We checked the survival and dissected the surviving oysters, after taking a haemolymph sample through the predrilled hole. We cut around  $25 mm^3$  ( $\sim 100 mg$  wet weight) of the mantle, gills and gut tissues with a sterile knife and flushed them thoroughly with sterile PBS in order to remove transient, non-attached bacteria. We immediately froze half of the tissue for microbiota analysis at  $-80^{\circ}C$ , while the other half was used to determine the number of cultivable *Vibrionaceae*. To do so, we homogenized the tissue pieces in  $500 \mu l$  of sterile PBS in the Qiagen TissueLyser (Hilden, Germany) using a single 5 mm stainless steel bead at 20 Hz for 3 min and plated  $10 \mu l$  on TCBS agar.

To determine the background composition of the microbial communities and the number of cultivable *Vibrionaceae* in the seawater, we took seawater samples (100 ml) on three occasions during the sampling period (six samples in total, three of them sequenced). The samples were filtered onto  $0.2 \mu m$  47 mm Nuclepore Track-Etch Membrane filters, which were then used for DNA extraction.

### Oyster immune parameters

In order to measure THC,  $50 \mu l$  of haemolymph was mixed with equal amounts of 6% formaldehyde in sterile seawater (SSW) and marine anticoagulant solution (Fedders and Leippe, 2008). The resulting solution was further diluted  $3 \times$  in sterile PBS and the cell count was measured with an automated cell counter (Scepter, Millipore, Darmstadt, Germany).

For phagocytosis, we followed the established protocols (Wendling and Wegner, 2013). In short,  $3 \times 60 \mu l$  of haemolymph were allowed to adhere to the bottom of 96-well plates for 1 h. The supernatant was carefully decanted, and the neutral-red-stained zymosan solution (Sigma-Aldrich) was added to haemocytes and incubated with shaking for 1 h. The reaction was stopped by the addition of 6% formol in SSW. The wells were washed several times with PBS, the haemocytes with the phagocytosed particles were solubilized in acidified ethanol (1% acetic acid, 50% ethanol), and the



absorbance was measured at 550 nm with Nanodrop ND-1000 spectrometer (peqlab, Erlangen, Germany). The standard curve was constructed from zymosan solution samples with known particle concentration, and the results were expressed as the number of phagocytosed particles per haemocyte.

To estimate the plasma protein content, 200  $\mu$ l of haemolymph was centrifuged for 5 min at 5000 *g*. The protein concentration in the supernatant was measured in triplicates with Quick-Start Bradford protein Bio-Assay (Bio-Rad, Hercules, CA, USA) according to the manufacturer's protocol.

#### DNA extraction

DNA was extracted from approximately 200  $\mu$ l of haemolymph, or approximately 50 mg of mantle, gill and gut tissue with Wizard SV 96 Genomic DNA Purification System (Promega, Mannheim, Germany). The samples were placed in pre-cooled ( $-20^{\circ}\text{C}$ ) TissueLyser Adapters (Qiagen) and homogenized in a mixer mill (Retsch, Haan, Germany), using a mixture of 0.5 mm glass-zirconium beads and a single 5 mm tungsten bead (for mantle, gill and gut) or 1 mm glass beads (for haemolymph) at 30 Hz for 5 min in order to completely disrupt the tissues and bacterial cells. The standard protocol for DNA extraction from animal tissues was used for mantle, gill and tissue, while only proteinase K (20  $\mu$ l of 20 mg  $\text{ml}^{-1}$  solution, Sigma-Aldrich) was added to the haemolymph for protein digestion. The samples were incubated at  $55^{\circ}\text{C}$  for at least 4 h and extracted according to the manufacturer's protocols. Blank extractions to check for bacterial contamination of reagents were also performed.

For the seawater samples, the filters were cut with sterile scissors into smaller pieces and homogenized in 2 ml tubes with a mixture of beads from PowerWater® DNA Isolation Kit (MO BIO Laboratories, Carlsbad, CA, USA) and 0.5 mm glass-zirconium beads in RLT buffer (DNeasy Blood & Tissue Kit, Qiagen) and further treated as described in Thomsen and colleagues (2012). Shortly, a round of bead beating at 30 Hz for 5 min was followed by 10 min at  $56^{\circ}\text{C}$  with continuous mixing. This was repeated twice and then the proteinase K was added and the digestion mix was incubated for 2 h. The samples were then extracted following the manufacturer's protocol with the adjusted reagent volumes. We used different methods for DNA extraction from the seawater and oyster tissues due to the differing properties of the source material. Although this could introduce a bias and caution is required in interpreting the results, previous research has shown that such biases tend to be minor (Sergeant *et al.*, 2012; Rubin *et al.*, 2014).

#### Polymerase chain reaction (PCR)

We amplified 16s rRNA V1–V2 regions with uniquely barcoded 27f and 338r PCR primers. The PCR reactions (25  $\mu$ l) were set up in 96-well plates as follows: 4  $\mu$ l of each forward and reverse primer (final concentration: 0.28  $\mu\text{M}$ ), 0.5  $\mu$ l dNTPs (final concentration: 200  $\mu\text{M}$  each), 0.25  $\mu$ l Phusion Hot Start II High-Fidelity DNA Polymerase (0.5 unit per reaction) and 5  $\mu$ l of High fidelity (HF) buffer (7.5 mM  $\text{MgCl}_2$ , Thermo Fisher Scientific, Waltham, MA, USA). We used 1  $\mu$ l of undiluted haemolymph DNA, 1  $\mu$ l of  $10 \times$  diluted

seawater DNA and 2–4  $\mu$ l of solid tissue DNA per reaction. For each 96-well plate, 20 control reactions (12.5  $\mu$ l) were performed: one positive control and unique combinations of all used forward and reverse primers as negative control, with water as a template.

The PCR cycling conditions were as follows: 30 s initial denaturation at  $98^{\circ}\text{C}$ , then 30 cycles: 9 s denaturation at  $98^{\circ}\text{C}$ , 1 min annealing at  $55^{\circ}\text{C}$ , 90 s extension at  $72^{\circ}\text{C}$ , 10 min final extension at  $72^{\circ}\text{C}$ .

In order to check for the product and to estimate its amount, the reactions were analysed immediately on a 1.5% agarose gel. Briefly, 5  $\mu$ l of loading buffer was mixed with 3  $\mu$ l reaction and loaded into the gel prepared with SYBR Safe DNA Gel Stain (Life Technologies GmbH, Darmstadt, Germany), including 3  $\mu$ l of O'GeneRuler™ 100 bp Plus DNA Ladder (Thermo Fisher Scientific). The separation was conducted at 120 V per 35 cm for 80 min. The result was photographed with Gel Doc™ XR+ System and analysed with Image Lab™ Software (Bio-Rad) to estimate the absolute concentration of the PCR products using the ruler as internal standard. If there was no amplification in blank extractions, they were excluded from further analysis. Equal amounts of the products from a single gel were pooled together (25–100 ng per sample), run on 1.5% agarose gel and purified with MinElute Gel Extraction Kit (Qiagen) according to the manufacturer's protocol, including optional additional centrifugation and steps recommended for salt-sensitive applications. The concentration of DNA in the resulting subpools was measured fluorometrically with Qubit dsDNA br Assay Kit (Life Technologies GmbH) in the Qubit fluorometer (Life Technologies Invitrogen GmbH, Karlsruhe, Germany). The equal amounts of subpools were then mixed together and frozen at  $-20^{\circ}\text{C}$  until sequencing. The paired-end sequencing by synthesis was performed on an Illumina MiSeq platform at the Max Planck Institute for Evolutionary Biology in Plön, Germany.

#### Sequence quality control and preprocessing

All sequencing libraries were processed together. Quality control, OTU clustering and taxonomy assignment were performed in MOTHUR (Schloss *et al.*, 2009), following the MOTHUR MiSeq SOP (Kozich *et al.*, 2013). We retained only overlapping regions of the contigs and removed any sequences with ambiguous bases and/or homopolymers of 8 bp or longer in order to ensure good quality and reduce the number of spurious OTUs. The sequences were aligned to SILVA 119 reference alignment (Quast *et al.*, 2013) cut to V1–V2 region, and the taxonomy was assigned with 80% confidence cut-off, using the Greengenes taxonomy 13\_08 (DeSantis *et al.*, 2006) and the Naïve Bayesian Classifier (Wang *et al.*, 2007) implemented in MOTHUR. Unknown (i.e. sequences not assigned to any kingdom), chloroplast, Archaea and Eukaryotic sequences were removed from further analysis. We performed single-linkage pre-clustering with two differences allowed (Huse *et al.*, 2010), removed the chimeras and created 97% OTUs using average-linkage clustering method. Consensus taxonomy for an OTU was assigned with a 50% consensus confidence threshold. We calculated rarefaction curves of diverse  $\alpha$ -diversity metrics in QIIME (Caporaso *et al.*, 2010) in order to estimate the effect of sampling effort and to determine sequencing depth for the

final analysis. We subsampled the dataset to 8000 reads per sample for the final analysis. Because the abundant OTUs (> 100 reads) in the single positive blank control were rare (< 1%) in the remaining samples, we simply excluded them from further analysis. In order to calculate a tree needed for phylogenetic measures of diversity, we picked a representative set of sequences using the distance method in MOTHUR and calculated the tree using FASTREE (Price *et al.*, 2010). The final dataset comprised 3 seawater, 141 haemolymph (including pre- and post-deployment samples), 68 gill, 61 gut and 62 mantle samples (335 in total).

### Statistical analysis

All statistical analyses were performed in R (R Core Team, 2013). For  $\alpha$ -diversity, we used a complete rarified dataset (8000 reads per sample) to analyse the differences in evenness [calculated as  $H/\ln(S)$ , where H is the Shannon–Wiener index and S is species richness] and species richness (total number of species). We first tested for differences between the seawater and oysters using non-parametric asymptotic Wilcoxon Mann–Whitney RS test (Wilcoxon RS test). We subsequently tested for differences between the tissues using robust analysis of variance (Wilcox and Schönbrodt, 2014), as the assumptions for classical ANOVA were not satisfied. Finally, we analysed haemolymph and solid tissues separately with linear mixed models (Bartoń, 2014; Kuznetsova *et al.*, 2015; Bates *et al.*, 2015).

Because not only *Vibrio* sp. but also other *Vibrionaceae* grow on TCBS agar and, in addition, classification of short reads to low taxonomic levels can be unreliable, we decided to focus the analyses on the whole *Vibrionaceae* family, and not just the genus *Vibrio*. In addition, due to the sampling procedure, the CFU counts were not directly comparable between the haemolymph and solid tissues, and were thus analysed separately.

For  $\beta$ -diversity, we kept only the OTUs with relative abundance higher than 0.1% in at least 10 samples to reduce the dataset complexity. We calculated Bray–Curtis distances and weighted UniFrac distances (Hamady *et al.*, 2010) using the *phyloseq* package (McMurdie and Holmes, 2013), and the results were further analysed by NMDS and Permanova (non-parametric permutational multivariate analysis of variance; Anderson and Beaven, 2001) implemented in the *adonis* function in the *vegan* package (Oksanen *et al.*, 2013). We first compared the tissues and then analysed the  $\beta$ -diversity in each tissue separately.

We statistically examined the variation at the class-level taxonomical composition between the tissues and the changes in the abundance of OTUs and genera in haemolymph microbiota according to origin and treatment by multivariate generalized mixed models (*mvabund* package; Wang *et al.*, 2012). This method fits a multivariate model for the differences between the whole communities as well as univariate models for the abundance of each taxon separately, thus identifying taxa responsible for the observed differences. Because the mean–variance relationship of the data is empirically estimated prior to the model-fitting (in this case negative-binomial), potential confounding of location and dispersion effects inherent to distance-based methods is avoided (Warton *et al.*, 2012).

We included time as an ordered factor in the models to check for temporal trends in the data. However, we could not disentangle individual variability from true time effects due to our experimental design. For this reason we do not further discuss the temporal trends, although we included them in results for completeness.

To explore both positive and negative associations of the OTUs within and between the tissues, we constructed an association network using the sparcc algorithm (Friedman and Alm, 2012) implemented in MOTHUR. We performed 10 000 permutations and kept only correlations > 0.4 with *P* value <  $10^{-4}$  to exclude as many spurious correlations as possible (Marino *et al.*, 2014). The input matrix was organized similar to Faust and colleagues (2012): OTUs in tissues in the rows and individual oysters in columns. Only the oysters with available data for all four tissues and the OTUs that appeared in at least one third of the samples were analysed (Berry and Widder, 2014). We statistically determined the significance of observed connectivity within/between the tissues by comparison to connectivity between the random subsets of nodes of equal size as the tested group (Faust *et al.*, 2012). Additionally, to assess the effect of treatment and origin on microbial associations in the haemolymph, we constructed a network for each experimental group of oysters including only haemolymph samples, and calculated their descriptive statistics including clustering coefficient (Newman *et al.*, 2002) and modularity. Modularity is a measure of separation of a network into densely intra-connected areas, with fewer connections between them (modules, communities), with respect to some division rule. In this case modules were computed using the walktrap algorithm (Pons and Latapy, 2005). The networks were visualized using the *igraph* package (Csardi and Nepusz, 2006). Raw demultiplexed sequence data are available at European Nucleotide Archive under the study accession number PRJEB8492.

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## Supporting information

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

**Fig. S1.** Correlation between the number of cultivable Vibrionaceae and Vibrionaceae evenness (A, B) or Vibrionaceae species richness (C, D) in the haemolymph and solid tissues. The lines show a linear model fit and are used only as an illustration for the relationship. All conclusions are based on non-parametric Spearman's  $\rho$  coefficient.  $\rho =$  Spearman's  $\rho$ .

**Fig. S2.** Association networks of haemolymph OTUs grouped by oyster origin and treatment showing only positive correlations (colour, shape and size legend as in Fig. 6). Separate subnetworks thus represent negatively correlated modules of the complete network. Arrows point to the 'sea-water' cluster, while the ellipses encircle all parts of the network where the other OTUs/motifs shared by the oyster groups are found.

**Table S1.** Pearson residuals for the  $\chi^2$  test of end-point survival between the four oyster groups. Texel non-antibiotic treated oysters are the only group with the higher than expected mortality.

**Table S2.** Multivariate generalized linear model (negative binomial) showing significant differences at the class level between the oyster tissues, origin and treatment.

**Table S3.** Multivariate generalized linear model (negative binomial) showing effects of oyster origin and treatment on the abundance of genera within the haemolymph. Only genera with significant differences are listed.

**Table S4.** Multivariate generalized linear model (negative binomial) showing effects of oyster origin and treatment on the OTU abundance within the haemolymph. Only OTUs with significant differences are listed.

**Table S5.** Properties of original haemolymph networks and average of 1000 random networks with same number of nodes and edges.

**Table S6.** Multivariate generalized linear model (negative binomial) showing effects of oyster origin and treatment on the genera abundance in solid tissues. Only genera with significant differences are listed.