Dual transcriptomics reveals co-evolutionary mechanisms of intestinal parasite infections in blue mussels *Mytilus edulis*

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Abstract
On theoretical grounds, antagonistic co-evolution between hosts and their parasites should be a widespread phenomenon but only received little empirical support so far. Consequently, the underlying molecular mechanisms and evolutionary steps remain elusive, especially in non-model systems. Here, we utilized the natural history of invasive parasites to document the molecular underpinnings of co-evolutionary trajectories. We applied a dual-species transcriptomics approach to experimental cross-infections of blue mussel *Mytilus edulis* hosts and their invasive parasitic copepods *Mytilicola intestinalis* from two invasion fronts in the Wadden Sea. We identified differentially regulated genes from an experimental infection contrast for hosts (infected vs. control) and a sympatry contrast (sympatric vs. allopatric combinations) for both hosts and parasites. The damage incurred by *Mytilicola* infection and the following immune response of the host were mainly reflected in cell division processes, wound healing, apoptosis and the production of reactive oxygen species (ROS). Furthermore, the functional coupling of host and parasite sympatry contrasts revealed the concerted regulation of chitin digestion by a Chitotriosidase 1 homolog in hosts with several cuticle proteins in the parasite. Together with the coupled regulation of ROS producers and antagonists, these genes represent candidates that mediate the different evolutionary trajectories within the parasite’s invasion. The host–parasite combination-specific coupling of these effector mechanisms suggests that underlying recognition mechanisms create specificity and local adaptation. In this way, our study demonstrates the use of invasive species’ natural history to elucidate molecular mechanisms of host–parasite co-evolution in the wild.

**KEYWORDS**
biological invasion, coevolution, de novo transcriptomics, host–parasite interactions, immune response, macroparasite

1 | INTRODUCTION

Molecular evolution is a comparatively slow process, and rapid adaptation requires either a large amount of standing genetic variation (Barrett & Schluter, 2008), or plastic mechanisms, for example, gene regulation or epigenetics (e.g., Rey, Danchin, Mirouze, Loot, & Blanchet, 2016; Shama et al., 2016). The interaction between hosts and parasites is a good example of rapid and

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dynamic evolution (Brockhurst, Morgan, Fenton, & Buckling, 2007; Buckling & Rainey, 2002; Decaestecker et al., 2007). Here, the host can evolve resistance to prevent a parasite from infecting, or evolve tolerance to minimize the damage by the parasite (Schneider & Ayres, 2008). The parasite, in turn, maximizes its fitness by evolving infectivity and virulence leading to the antagonistic arms race between host and parasite, which is characterized by rapid reciprocal adaptation (e.g., Bishop, Dean, & Mitchell-Olids, 2000). Although co-evolution is a key assumption of host-parasite evolutionary theory, empirical support is lacking far behind with only few studies supporting predictions of co-evolution. Most of these originate from the laboratory (e.g., Berenos, Wegner, & Schmid-Hempe, 2011; Buckling & Rainey, 2002; Eizaguirre, Lenz, Kalbe, & Milinski, 2012) and very few come from the wild (Decaestecker et al., 2007). The phenotypes of characterizing these antagonistic interactions may appear similar, but the underlying molecular mechanism may differ substantially. Conversely, seemingly different phenotypic trajectories may also be based on the same molecular machinery that merely differs in its expression levels (Elmer et al., 2014; Hanson, Hu, Hendry, & Barrett, 2017). Overall, our mechanistic understanding, and consequently also our understanding of the mode and speed of co-evolution, is so far limited.

Parasite invasions are ideal to study the temporal dynamics of host-parasite evolutionary interactions in the wild (Goedknegt et al., 2015). For instance, it is possible to use the implicit space for time substitution (Reusch, 2014) of invasive parasites along their invasion route to study host-parasite co-evolution along a time axis covering ancient to no sympatry (Feis et al., 2016). The invasion routes of the parasitic copepod Mytilicola intestinalis into the Wadden Sea unites all aspects of an invasion to study host-parasite co-evolution (Feis et al., 2016). Native in the Mediterranean and Adriatic Seas, M. intestinalis invaded mussel beds in the late 1930s (Caspers, 1939) and has spread southwest towards the island of Texel (The Netherlands) (Drinkwaard, 1993) and northward to Sylt (Germany) (Dethlefsen, 1972), arriving at both islands around the same time in the early 1970s. The invasion routes are thus replicates of a newly begun co-evolutionary relationship between invasive blue mussel Mytilus edulis and the parasite M. intestinalis. Although the parastic lifestyle of M. intestinalis was sometimes debated (Davey, Gee, Bayne, & Moore, 1977), we could already show that M. intestinalis and mussels have evolved different evolutionary trajectories along their invasion routes. Specifically, sympatric combinations of M. intestinalis and mussel hosts caused more damage to the host than allopatric combinations (Feis et al., 2016), probably owing to fact that Mytilicola parasites mainly consume host tissue (Goedknegt et al., 2017). Mussels from the southwest invasion front on Texel, for example, were more resistant than those from Sylt, while mussels from Sylt tended to be more tolerant to being parasitized (Feis et al., 2016).

To understand the evolutionary process leading to the phenotypic differences, the molecular mechanisms in the mussel–Mytilicola relationship need to be deciphered. The major immune pathways are reasonably well characterized in bivalves and gastropods (Gerdo, 2017). For mussels (Mytilus spp.), these were mainly derived from transcriptional and metabolite studies in response to microbial pathogens, for example, Gram-positive and Gram-negative bacteria (Liu et al., 2014; Toubiana et al., 2014; Venier et al., 2011), yeast and filamentous fungi (Sonthi et al., 2012). Because -omics tools were essential in the description of important pathways in mussel immunity (Gerdo & Venier, 2015), this approach can also be useful to characterize the immune response towards macroparasitic infections. Macroparasites have been targeted only by very few mussel studies (e.g., trematode infections, Gorbushin & Lakovleva, 2011), despite a high parasite richness of this widespread epibenthic filter feeder (Thieltges, Engelsma, Wendling, & Wegner, 2013). Methodologically, macroparasite infections have the advantage that host and parasite transcriptomes can be sequenced separately, extending a dual transcriptomics approach to a dual-species transcriptomics approach. In microbial infections, hosts and parasites often cannot be separated physically and classical dual transcriptomics relies on bioinformatic separation of host and parasite transcripts leading to heavy bias towards the host with only few parasite transcripts (Greenwood, Ezquerra, Behrens, Branca, & Mallet, 2016). Nevertheless, a dual transcriptomics approach can be very helpful to identify interacting genes (Schulze, Schleicher, Guthke, & Linde, 2016; Westermann, Gorski, & Vogel, 2012) and has been applied to several model organisms (e.g., Choi, Aliota, Mayhew, Erickson, & Christophersen, 2014; Pittman, Aliota, & Knoll, 2014; Rosani et al., 2015; Tierney et al., 2012). For bivalves, the interaction between the Pacific oyster Magallana gigas (previously known as Crassostrea gigas, see Salvi & Mariottini, 2017) and Ostreid Herpesvirus type 1 (OShV-1) has been characterized using this approach, revealing that OShV-1
interfered with active host defence by suppression of cytokine signalling through interferon-related pathways (Rosani et al., 2015).

So far, these dual transcriptomics studies have mainly focused on the infection process as such, but have not incorporated the geographic aspects of co-evolutionary rapid local adaptation or contrasted sympatric and allopatric combinations of hosts and parasites.

Here, we incorporate these aspects by combining the natural history of the Mytilicola invasion with a dual-species RNA seq analysis of a reciprocal infection experiment. This approach can uncover the molecular underpinnings of the mussel-Mytilicola relationship at the transcriptional level and thus help explain the different evolutionary trajectories observed along the invasion route of Mytilicola (Feis et al., 2016). We characterize this interaction on three different levels representing our explicit experimental contrasts. First, we ask which genes and processes are involved in regulatory responses of hosts infected by Mytilicola by contrasting experimentally infected hosts with control hosts, representing the infection contrast. Second, we identify host genes that are differentially regulated in infections with locally co-evolved parasites compared to infections with parasites that lack a recent shared co-evolutionary history, representing the sympatry-allopatry contrast. Lastly, our dual-species transcriptomics approach allows us to use the same sympatry-allocapty contrast to identify differentially regulated genes in the parasite, which can be matched to the host sympatry-allocapty contrast, and couple phenotypic evolutionary trajectories to co-evolved gene regulatory processes in antagonistically interacting species.

2 | MATERIALS AND METHODS

2.1 | Experimental procedures

The details of the underlying cross-infection experiment are described in Feis et al. (2016). Briefly, mussels M. edulis and parasitic copepods M. intestinalis were collected at two locations, representing the two invasion pathways of the parasite. These were at Vlakte van Kerken on Texel, The Netherlands, representing the southwestern pathway and in Königshafen on Sylt, Germany, representing the northern pathway. The mussels used in the experiment were treated against any previous Mytilicola spp. infections according to the method of Blateau, Le Coguic, Mailhe, and Grizel (1992) and were left to recover after the anti-Mytilicola treatment for at least 2 weeks before the experiment started. The copepods used in the experiment to infect mussels were hatched from egg sacks of gravid M. intestinalis from the two locations. Mussels were infected with 24 copepodes from one or two mothers. Therefore, copepodes within one infection were related; however, we do not know whether multiple males sired the offspring from one female. Assuming that relatedness within the population will be 0, we can be sure that relatedness was higher within a single infection than between infections. In a full-factorial design, we generated different experimental contrasts, that is, infected vs. uninfected control mussels, sympatric (Texel host × Texel parasite, Sylt host × Sylt parasite) and allopatric (Texel host × Sylt parasite and vice versa) host and parasite combinations. In the control treatment, we did not expose mussels to any copepods. Each mussel was kept in a temperature-controlled room at 18°C in its own bottle that was supplied with its own flow-through inlet distributing filtered seawater equally (salinity of the water input was 29–30 psu). Bottles with mussels were randomly relocated within and between containers twice per week. Mussels were dissected under a stereomicroscope 80 days after the experiment had started. The intestinal tissue of each mussel, which is in direct contact with the parasite, and M. intestinalis specimens were put in RNA later (Qiagen, Hilden, Germany) separately and were frozen at −80°C until further processing.

2.2 | RNA extraction, library preparation and sequencing

RNA extractions of mussels and parasites were done with the RNeasy Plant Mini Kit (Qiagen, Hilden, Germany). Since extraction of single female parasites did not yield enough RNA for sequencing, we pooled three female M. intestinalis per sample. We had three biological replicates for each treatment combination (see Table 1), except for the Texel host × Texel parasite combination, for which we had two biological replicates as not enough material was present for pooling a third replicate. Total RNA concentrations were measured on a NanoDrop ND-1000 spectrophotometer (PepLab [VWR], Erlangen, Germany). The concentration, purity and integrity of the RNA were further measured and checked with an Agilent 2100 Bioanalyzer using the Agilent RNA 6000 Nano Kit (Agilent Technologies, Waldbronn, Germany). Library preparation was done with the TruSeq Stranded mRNA HT Library Prep Kit (Illumina, San Diego, CA, USA). We used an Agilent 2100 Bioanalyzer with the Agilent 7500 Nano Kit (Agilent Technologies, Waldbronn, Germany) to quantify the amount of cDNA in each library and to check the quality of each cDNA library, after which the cDNA libraries were pooled equimolarly. All of the above kits were used according to the manufacturer’s protocols. Next-generation sequencing was performed on a NextSeq500 DNA sequencer (Illumina, San Diego, CA, USA) at the Alfred Wegener Institute, Helmholtz Centre for Polar and Marine Research with a single-end sequencing strategy at a read length of 75 bp. Illumina BCL files were converted to fastq files and demultiplexed using bcl2fastq (v2.17, Illumina, San Diego, CA, USA) with default settings. Trimmomatic (Bolger, Lohse, & Usadel, 2014) was used to clean raw reads by filtering out short (<36 bp) and low-quality reads (sliding window option), as well as adapters if still present.

2.3 | Mytilus edulis: assembly and annotation

Reads from 18 M. edulis gut libraries (BioSample Accession nos. SAMN08395021-SAMN08395038 in BioProject PRJNA430138) presented here were combined with reads from a further 27 M. edulis gut libraries (BioSample Accession nos. SAMN08395039-SAMN08395065 in Bioproject PRJNA430138) for de novo assembly of a mussel transcriptome backbone. All libraries were derived from the same experiment with the same protocols. We chose to make
transcriptome backbones with all libraries for assembly robustness and analysed only the first subset of 18 *M. edulis* libraries that were relevant for the current question. The backbone was assembled de novo in CLC Genomics Workbench 8.5.1 (CLCbio, Denmark) with default parameters, using automatic word and bubble size and a minimum contig size of 200 nucleotides. Contigs were updated by back-mapping of reads (mismatch cost 2, insertion and deletion cost 3, length fraction 0.5 and similarity fraction 0.8). The mussel transcriptome backbone was annotated using the Trinotate pipeline (Grabherr et al., 2011), including BLAST (Altschul, Gish, Miller, Myers, & Lipman, 1990) and Gene Ontology (GO) annotation (The Gene Ontology Consortium, 2000).

### 2.4 *Mytilicola intestinalis*: assembly and annotation

Reads from 11 *M. intestinalis* libraries (BioSample Accession nos. SAMN08394998-SAMN08394998 in BioProject PRJNA430138) presented here were combined with reads from 13 *M. intestinalis* libraries (BioSample Accession nos. SAMN08394999-SAMN08395006, SAMN08395008 and SAMN08395010 in BioProject PRJNA430138) from another experiment for de novo assembly of a parasite transcriptome backbone in CLC Genomics Workbench 8.5.1 (CLCbio, Denmark) with identical parameters used for the mussel assembly (see above). All libraries were derived from the same experiment with the same protocols, but only the 11 *M. intestinalis* libraries presented here were relevant for the question in this manuscript. The parasite transcriptome backbone was annotated by BLAST of the contig sequences against UniRef, only retaining hits with an e-value cut-off of <1 x 10^-5. We used TransDecoder to find the most probable ORFs (Haas et al., 2013) and InterProScan (Jones et al., 2014) to gain more insight into the potential function of differentially expressed *M. intestinalis* contigs without a BLAST annotation. After removing contigs that had a bivalve (host) or phytoplankton (food) annotation, 44,616 contigs remained (Table 2 for assembly statistics). The quality of the parasite transcriptome backbone was assessed through BUSCO (Simao, Waterhouse, Ioannidis, Kriventseva, & Zdobnov, 2015) (Table S1).

The same gene in different organisms may have different functions, but belong to the same gene regulatory network (e.g., Hinman & Davidson, 2007). Understanding the potential pitfalls of inferring function from annotations based on sequence similarity without further experimental evidence on each specific gene for mussel and *M. intestinalis*, we will assume homologous function for the differentially expressed genes we report.

### 2.5 Differential expression analyses

Differential expression analyses and GO enrichment analysis were performed in the R statistical environment version 3.4.1 (R Core Team 2017), after the reads from each library were mapped back to the respective transcriptome backbone, using the RNA seq tool in CLC Genomics Workbench 8.5.1 (CLCbio, Denmark), with the length and similarity fraction parameters set to 0.5 and 0.8, respectively. The resulting count data were further analysed using the DESeq2 package (Love, Huber, & Anders, 2014) from BioConductor (Huber et al., 2015), where we selected the 18 transcriptomes of mussels and 11 of parasites that belong to the experimental groups described in this article.

To address the different questions for mussel hosts, we first analysed the complete mussel RNAseq data set to examine the difference between all infected vs. all control mussels and then focused on a subset including only the infected mussels to differentiate between sympatric and allopatric infections. For the parasite, we only contrasted sympatric and allopatric infections. The DESeq experiment design was full rank for all data sets. To include only meaningfully expressed contigs, we applied a mapping cut-off filter by excluding contigs with fewer than five hits per sample on average. Main effects and

### TABLE 1 Overview of the number of replicates successfully sequenced for each experimental treatment. For the parasite RNAseq data, the first number represents the number of biological replicates and the second number the amount of parasites pooled within each replicate

|        | Host RNAseq data | Parasite RNAseq data |
|--------|------------------|-----------------------|
| Texel mussel | Mytilicola intestinalis | Sylt M. intestinalis No parasite |
| 3 | 3 | 3 |
| 2 x 3 | 3 x 3 |
| Sylt mussel | 3 | 3 |
| 3 | 3 |

### TABLE 2 Summary of sequencing, assembly and annotation statistics

|        | Mytilus edulis | Mytilicola intestinalis |
|--------|----------------|-------------------------|
| Sequencing statistics | | |
| Number of clean reads | 347,996,951 | 108,116,930 |
| Total bases | 26,215,385,329 | 8,144,066,749 |
| % GC content | 40.8% | 50.2% |
| Assembly statistics | | |
| Number of contigs | 163,270 | 45,645 |
| Average contig length | 641 | 738 |
| N50 contig length | 1,206 | 1,289 |
| Range of contig lengths | 200–13,745 | 200–19,927 |
| Annotation statistics | | |
| Annotated contigs by blastx | 29,799 (18.3%) | 12,653 (27.8%) |
| Annotated contigs by blastp | 22,451 (13.8%) | 16,259 (35.7%) |
| Annotated contigs with GO | 28,764 (17.6%) | 12,826 (28.1%) |
| TOTAL annotated contigs | 30,514 (18.7%) | 16,259 (35.7%) |

N50 = 50% of the bases are in contigs larger than this number; GO, Gene Ontology.
interaction effects were tested by Wald tests. For each main and interaction effect, we used the fdrtool for false discovery rate correction with the Benjamin–Hochberg method (Strimmer, 2008a, 2008b). We consider contigs differentially expressed if the adjusted p-value \( p_{\text{adj}} < \cdot05 \) and the absolute log2fold change (LFC) \( \geq 1.5 \).

To gain further insights into the functions of differentially expressed contigs for mussels, we performed GO enrichment analyses for biological processes of the main and interaction terms based on the GO annotations from Trinotate using the R-package topGO (Alexa, Rahnenführer, & Lengauer, 2006). A similar approach for *M. intestinalis* was not successful because of the low number of differentially expressed contigs with GO annotations. Due to their functional relevance in the interaction with *M. intestinalis*, we examined the differentially expressed mussel genes with the GO annotations (including offspring) in the biological processes of "response to stimulus" (GO:0009611), "immune system processes" (GO:0002376) and "death" (GO:00016265) closer, and in addition, we manually curated this DEG set by including contigs with the same or similar protein name that lacked a GO annotation.

### Results

#### 3.1 Mussel and *Mytilicola* transcriptome backbone quality

We obtained a total of 348 million cleaned reads for the mussel *M. edulis* and 108 million cleaned reads for the parasite *M. intestinalis*, resulting in assemblies of 163,270 and 45,645 contigs, respectively (Table 2). Applying our mapping cut-off filter resulted in 63,577 contigs for the infection vs. control differential expression analyses and in 64,759 mussel and 19,546 parasite contigs for the sympatry vs. allopatry contrast.

#### 3.2 Differential expression analysis in mussels: infection vs. control

We found a total of 471 differentially expressed contigs \( p_{\text{adj}} < \cdot05 \) and LFC \( \geq 1.5 \). 233 had an annotation, Table S2: 226 in the infection contrast, 176 in the host source contrast and 184 for the interaction infection by host source (Figure 1a, Table S2). Of those genes, 116, 30 and 17 in the respective contrasts belonged to significantly enriched biological processes identified by GO enrichment analyses. The most represented GO terms enriched by infection with *Mytilicola* belonged to cellular process (111 genes, GO:0009987, Fisher’s exact test, \( p = .0286 \)), metabolic process (90 genes, GO:0008152, Fisher’s exact test, \( p = .0125 \)), single-organism process (100 genes, GO:0044669, Fisher’s exact test, n.s.) and biological regulation (57 genes, GO:0065007, Fisher’s exact test, n.s.). Significantly enriched offspring terms of cellular process and single-organism process were mitotic cell cycle (GO:0000278, Fisher’s exact test, \( p = 9.3e-06 \)) and cell division (GO:0051301, Fisher’s exact test, \( p = .00023 \)). The GO process cellular component of organization or biogenesis (GO:0071840) was only found in the infection contrast. Its significant offspring terms included mitotic nuclear division (GO:0007067, Fisher’s exact test, \( p = 6.33e-06 \)), ribosome biogenesis (GO:0042254, Fisher’s exact test, \( p = .0116 \)) and mitotic spindle assembly (GO:0090307, Fisher’s exact test, \( p = .000987 \)). Another significantly enriched term in the infection treatment was wound healing (GO:0042060, offspring of response to stimulus, Fisher’s exact test, \( p = .0166 \)), suggesting that tissue damage caused by the parasite induced a tissue regeneration response. Offspring of the term reproductive process (GO:0022414) was significantly enriched in the infection contrast (12 genes) and in the interaction contrast (three genes), suggesting a resource shift from reproduction to wound healing when infected. Furthermore, the GO terms reactive oxygen species (ROS) metabolic process (GO:0072593) and response to oxidative stress (GO:0006979) were significantly enriched in the infection contrast (Fisher’s exact tests, \( p = 3.2e-05 \) and \( .00596 \), respectively).

Additionally, exploration of the data set including only the genes involved in immune system processes (GO:0002376), response to stimulus (GO:00050896) and death processes (GO:0016265) resulted in a list of 46 differentially regulated genes (Figure 2a). Most notably, infection led to an upregulation of genes inducing ROS, such as four dual oxidase homologs (LFC ranging 4.4–4.8), as well as ROS antagonists, such as homologs of glutathione peroxidase 1 and 2 (LFC = 1.7 and 2.4, respectively) and glutathione reductase (LFC = 2.3). This is a further indication that ROS pathways are among the most prominent immune mechanisms against *M. intestinalis* infections. As a possible consequence of increased ROS damage, we also found upregulation of genes involved in apoptosis, including stimulators such as homologs of inhibitor of growth protein 3 (LFC = 1.6) and peptidyl-prolyl cis-trans isomerase D (LFC = 2.5 and 2.6); these two contigs are probably fragments of the same transcript, because the protein sequences did not overlap upon alignment, data not shown), as well as homologs of the inhibitors filamin A (LFC = 1.7 and 1.6; these two contigs showed overlap in their alignment, data not shown), KIF14 (LFC = 2.1), and baculoviral IAP repeat-containing protein (LFC = 3.3). Some of the immunity-related genes were downregulated in infected mussels (Figure 2a). The downregulation of a homolog of Toll-like receptor 7 (TLR7, LFC = –3.2), a key component of innate immunity which recognizes molecular patterns specific to microorganisms, might indicate a shift of the immune response away from the Toll pathway (Toubiana et al., 2014). A homolog of sphingosine kinase 2 (LFC = –2.4), which may have pro- and anti-inflammatory functions (Neubauer & Pitson, 2013), was also downregulated in *M. intestinalis*-infested mussels.

#### 3.3 Differential expression analysis in mussels: sympatry vs. allopatry

In comparison with the infection contrast, we found more than twice as many differentially expressed contigs (946 at \( p_{\text{adj}} < \cdot05 \) and LFC \( \geq 1.5 \), of which 262 were annotated, Table S3). The majority (662 genes, 70%) of these were caused by the different parasite sources, while only 224 (24%) were differentially expressed between the host sources. Host by parasite combinations led to specific induction patterns in relatively few genes (165 genes, 17%, Figure 1b), indicating that the combination-
Specific phenotypic patterns (Feis et al., 2016) rely on the regulation of relatively few genes. Within these respective contrasts, 27, 5 and 21 genes belonged to significantly enriched biological processes in the GO enrichment analyses of the contrasts among different parasite sources, different host sources and host by parasite combinations, respectively. The term immune system process (GO:0002376) was not significantly enriched (nor any of its offspring terms, Figure 1b) in the parasite contrast GO enrichment analyses, but it was in both host contrast and host-parasite contrast (Fisher’s exact test, \( p = .00985 \) and \( p = .0436 \), respectively). Other significantly enriched GO terms in the host-parasite contrast were defence response (GO:0006952, \( p = .0225 \)), response to other organism (GO:0051707, \( p = .0128 \)) and response to bacterium (GO:0009617, \( p = .0037 \)), all of which are offspring terms of response to stimulus (GO:00050896).

The important role of parasite origin for host transcription becomes even more obvious in the curated data set that only contained genes involved in immune system (GO:0002376), response to stimulus (GO:0050896) and death processes (GO:0016265). Infection with the parasites from different sources changed regulation patterns in 30 out of 37 genes (81%), many of which were either stimulators or antagonists of ROS production (13 differentially expressed contigs: contig numbers 23,071, 41,162, 50,761, 5,027, 106,242 and 14,158 for ROS and contig numbers 1,728, 38,961, 14,694, 116,190, 90,969, 61,914, 105,058 for ROS antagonists). Regulation of these genes was not uniform, but varied depending on the parasite origin. A superoxide dismutase [Cu-Zn] homolog (SOD, a ROS antagonist) was upregulated (LFC = 15.5) only in the mussels infected with Sylt parasites, while homologs of chorion peroxidase,

![FIGURE 1](https://wileyonlinelibrary.com)
peroxidase-like protein, peroxidin (all antagonistic to ROS) and a dual oxidase homolog stimulating ROS production were all down-regulated (LFC = −20.5, −21.2, −2.7, −6.7, respectively). Similarly, several caspase 3 apoptosis homologs showed parasite-specific induction (contig numbers 126,399, LFC = 24.2; 47,524, LFC = 20.2; and 73,225, LFC = −19.6), suggesting that the regulation of genes with similar function can fine-tune functional responses at the phenotypic level. For the host contrast, only four genes (11%) were differentially regulated in the curated data set, most notably containing two Chitotriosidase 1 (Chit1) homologs that were less expressed in Sylt hosts (Figure 4). Chit1 is an enzyme that breaks down chitin. Both Chit1 contigs probably belong to the same gene, because no overlap existed between the aligned sequences (data not shown). Among the contigs significant for the host × parasite contrast (i.e., sympathy contrast, the interaction term in the model), we found nine genes with combination-specific regulation patterns (Figure S2). These included the apoptosis factor homologs of C1q TFN3, Toll-like receptor 6 (TLR6), Caspase 3 and Fat-like cadherin-related tumor suppressor (Figure 2b, LFC = 6.7, −3.17, −25.8 and −23.5, respectively). Apart from several dual oxidase homologs (contig numbers 23,071, 41,162, 50,761, 5,027 and 14,158) that stimulate ROS production and reacted to infection in general (Figure 1a; these contigs all showed LFC ≥ 1.5). In contrast to the host sympatry contrast, where most parasite combinations yielded 16 genes (30%), about half (26 genes) of the 53 differentially expressed genes had a significant BLAST E-value, suggesting that the regulation of genes with similar function can fine-tune functional responses at the phenotypic level.

3.4 Differential expression analysis in Mytilicola parasites

We found 53 differentially expressed contigs (p-value < .05 and LFC ≥ 1.5). In contrast to the host sympathy contrast, where most genes were differentially regulated depending on parasite source, in the parasite transcriptomes most differentially regulated genes were found between host sources (30 genes, 57%), while only 13 (25%) originated from differences between the parasites themselves. The interaction term representing regulation patterns for specific host × parasite combinations yielded 16 genes (30%). About half (26 genes) of the 53 differentially expressed genes had a significant BLAST or InterProScan hit, leaving 27 genes without putative functional annotation (Figure 3, Table S4).
Differences between the host sources were mainly due to down-regulation of genes in the parasites infecting Sylt hosts (Figure 3). Where homologies could be determined, these mainly included genes involved in development of the cuticle (four cuticle proteins, one chitin-binding domain, LFC range –1.8 to –4.4; Figure S1), which could indicate that the external physical barrier of the parasite suffered less damage in the guts of Sylt mussels—a pattern matching the downregulation of Chitotriosidase 1 in Sylt mussels (Figures 2, 4 and S2).

Out of the 13 differentially regulated genes between parasite sources, only three contigs downregulated in Sylt parasites had annotations: nephrin (contig number 3,349, LFC = –2.3), which plays a role in the development and function of kidneys in mice, aldehyde dehydrogenase (contig number 11,166, LFC = –2.3), which oxidizes aldehydes and may be involved in lipid peroxidation, and cathepsin K (contig number 8,083, LFC = –2.0), a protease with a high specificity for kinins (Figure 3). These genes are involved in catalytic processes and suggest differences in breakdown processes of complex molecules between the parasite sources.

Combination-specific regulation patterns in parasite genes led to upregulation of nine genes in sympatric combinations. These included homologs of Tick legumain (LFC = 1.8; Figure S1) putatively involved in the digestion of host macromolecules in other parasites, Kelch-like protein 10 (LFC = 1.7), involved in fertility, as well as a C-type lectin-like (LFC = 2.0), involved in sugar binding/immune recognition. As a direct link to host gene regulation, ROS antagonists (nicotinic oxide dioxygenase, LFC = –1.8; animal haem peroxidase, LFC = –4.0) of parasites were specifically downregulated in sympatric combinations (Figures 3 and S1), which matches the parasite-specific regulation of nicotinic oxide synthase in the host and might contribute to the lower condition of host in sympatric infections (Figure 2) (Feis et al., 2016). Parasites in sympatric combinations also seemed to allocate fewer resources to reproduction as indicated by the significant downregulation of two vitellogenin 2 homologs (Figures 3 and S1, LFC range –9.0 to –11.4).

4 | DISCUSSION

Our dual-species transcriptomic analyses clearly showed that the transcriptional response of both hosts and parasites was mainly driven by the differences between the respective antagonists. The parasite’s transcription mainly reacted to host origin, while host transcription reacted to the parasite origin. The resulting transcriptional response may thus be interpreted as a clear sign of co-evolutionary adjustment of gene expression in this host-parasite interaction. While our analysis only captures a single time point during the interaction of mussels and Mytilicola, we are confident that the chronic state of infection represents a relevant time point that integrates the interaction between host and parasite over an extended period. Adult female M. intestinalis cause the highest energy drain and can be found throughout the year in the field (F. Demann, unpublished data). In this way, we probably captured the chronic effects of the parasite rather than the acute ones, but such chronic effects can also be expected for the natural situation. At the functional level, we found that with 13 out of 37 differentially expressed annotated immune genes ROS pathways were important in the interaction between M. intestinalis and its mussel host. Because the ROS response is a fairly general and unspecific response to stress and diseases (Naviaux, 2012), there have to be gene regulatory mechanisms that create the observed specificity reflecting local adaptation by coupling of the matching host and parasite genes. Here, the higher expression of ROS-producing genes in sympatric hosts was coupled to ROS antagonist production in the corresponding parasites demonstrating this specificity (Figures 4, S1, and S2). This link between host and parasite transcriptomes was also reflected in other functionally related interactions. For example, chitin digestion of the hosts was coupled to cuticle protein expression in the parasite and might be a core mechanism underlying the resistance phenotype differences previously observed along the invasion fronts (Feis et al., 2016). Such coupled gene expression pattern not only demonstrates the usefulness of dual transcriptomics approaches (Greenwood et al., 2016), but, more importantly, it shows that regulation of functionally important differences evolved rapidly along the two invasion fronts, supporting the usefulness of invasive species’ natural history to understand host–parasite co-evolution in the wild.

4.1 | Mussel’s response to infestation by M. intestinalis

Mussels that were experimentally infected with M. intestinalis parasites regulated the expression of genes with functions in cell cycling and cell division and associated with wound healing (Figure 1a). *Mytilicola intestinalis* is known to obstruct the mussel’s intestine and perforate the intestinal wall with its appendages and hooks (Figueras, Jardon, & Cladras, 1991; Moore, Lowe, & Gee, 1977; Robledo, Santarém, & Figueras, 1994), and this wounding is reflected in the transcriptome of the host. Increased cell division suggests that wounds in the gut wall are being closed and the intestinal epithelium is being replaced after disruption to guarantee proper function of the gut. In addition, downregulation of anticoagulation agents (e.g., a homolog of fibrinolytic enzyme isozyme C) in *M. intestinalis*-infested mussels indicates an increased rate of blood clotting in infected mussels. Wounding and repair of physical damage therefore represent the major regulators of host gene expression.

Apart from the primary response to physical damage, host transcriptomes in the sympathy-allopatry contrast were also enriched in genes linked to an immune response. Similar to other studies investigating immune responses in mussels (e.g., Liu et al., 2014), we found the upregulation of genes involved in the production of ROS in *M. intestinalis*-infested mussels. This was accompanied by upregulation of genes providing antioxidants, likely to limit the negative effects of ROS on the host’s own tissues. This mechanism of defence is indeed general and does not only combat pathogen infections, but also plays a role in defence against infection by
We further found a differential regulation of the homolog maternal embryonic leucine zipper kinase, an enzyme that can be involved in apoptotic processes (Jiang & Zhang, 2013). Apoptosis may in part be a consequence of ROS production (Matès & Sánchez-Jiménez, 2000). Apoptotic processes play an important role in the immune reaction to foreign substances and were predicted to be important as first line of defence in the digestive tissues of mussels to macroparasites such as *M. intestinalis* (Romero, Novoa, & Figueras, 2015). Consequently, apart from the direct response to wounding, the upregulation of genes associated with cell division processes may also be linked to apoptosis by replacing apoptotic cells. Other immune genes that were differentially expressed in infected mussels (Figure 2a) were either annotated by host–virus interaction according to the UniProt database (Foulger et al., 2015; Masson et al., 2014; The UniProt Consortium, 2017), or were related to response to microorganisms. Intestinal parasites have indeed been shown to alter microbial gut homeostasis (e.g., Berrilli, Di Cave, Cavallero, & D’Amelio, 2012; Cariveau, Elijah Powell, Koch, Winfree, & Moran, 2014; Kreisinger, Bastien, Hauffe, Marchesi, & Perkins, 2015). The
downregulation of TLR7 in infected hosts may suggest a shift away from bacterial defence pathways (such as the Toll pathway) in *M. intestinalis*-infested mussels. Combined with the constant wounding of the gut epithelium, this might lead to a higher susceptibility towards secondary infections.

Chronic infection by *M. intestinalis* therefore orchestrates a multilayered cascade of transcriptional reactions in the mussel host, involving ROS mediated and apoptotic immune responses, both potentially damaging to host tissues. Together with the physical damage caused by lesions in the gut wall, these may lead to substantial wounding and cell division response to maintain gut homeostasis.

### 4.2 Dual transcriptomics: coupled gene expression with links to phenotype

The putative function of differentially regulated genes in the parasite was associated with metabolic processes such as proteolytic digestion. Here, especially the combination-specific expression pattern of the tick legumain homolog (contig number 1,525) could be a marker...
for parasite feeding activity on host tissue. Legumain is expressed in the midgut in blood digesting ticks, where it serves as a haemoglobinase, processing haemoglobin and activating other proteases (Dall & Brandstetter, 2016; Sojka, Francischetti, Calva, & Kotsyfakis, 2011). In other invertebrate groups (Fuzita et al., 2015 and references therein), but also in the protochordate Branchiostoma belcheri (Teng, Wada, & Zhang, 2009), legumain plays a role in food digestion: it breaks down macromolecules in the acidic gut environment. Thus, the higher legumain expression levels in sympatric combinations could be a direct indicator of lower host condition (Feis et al., 2016). Furthermore, other genes potentially involved in feeding (e.g., MAM and a low-density lipoprotein [LDL]-receptor class A [LDLRA] domain-containing protein) showed similar host–parasite combination-specific expression patterns, indicating once more a stronger exploitation of sympatric hosts. Arthropods have to acquire cholesterol from their diet as they are incapable of its de novo biosynthesis (Hassett, 2004; Perner et al., 2016). LDL is a rich source of cholesterol (Perner et al., 2016). Consequently, LDL receptors are also upregulated in scorpion digestion (Fuzita et al., 2015). The increased transcription of these genes in sympatric combinations might therefore indicate increased digestion of host tissue by Mytilicola, leading to higher energy drain from the host, which could further contribute to lower body condition in sympatric combinations (Feis et al., 2016).

Other genes with host–parasite combination-specific expression patterns in the sympatry contrast were involved in the parasite's immune response or defence against the host's immune response. The upregulation of a C-type lectin homolog in sympatric combination (Figure S1) suggests an increased immune activity of the parasite itself. C-type lectins are widespread pattern recognition molecules that are involved in carbohydrate recognition often associated with bacterial lipopolysaccharide in crustaceans (Kawabata & Iwanaga, 1999), suggesting that the parasite's immunological surveillance may be fine-tuned towards bacteria encountered in sympatric host guts. Since immune resources of the host are diverted to combat the parasite, less immune resources might be available to maintain bacterial homeostasis in the gut. Disturbed host microbiota can contain higher number of secondary opportunists (Lokmer & Wegner, 2015) that could potentially harm the parasite. This effect might be particularly strong in the already weakened sympatric hosts therefore calling for an increased level of immunological surveillance on the parasite's side as well.

A strong relative increase in host ROS-producer transcription after infection (Figure 4) was one of the main defence mechanisms. Here, the pattern of host ROS-producer transcription supports the hypothesis of a higher degree of self-damage in sympatric combinations (Feis et al., 2016). Host expression patterns of ROS antagonists, on the other hand, displayed opposite patterns, with especially high levels of host ROS antagonist expression in Texel hosts × Texel parasites combinations (Figure 4). However, the parasite expression profiles revealed that the parasites in Texel host × Texel parasite combinations hardly express any ROS antagonists. Assuming that transcription will positively correlate with ROS production this may indicate that the total amount of ROS scavengers might be lower and more ROS is freely available. A qualitatively similar pattern could be derived for the combination of Sylt hosts × Sylt parasites, where the increase in host ROS production expression was not associated with a remarkable increase in either host or parasite ROS antagonist expression, which might also result in a higher relative amount of freely available ROS in the mussel gut. The high level of ROS antagonist transcription in Sylt parasites in general might therefore explain the small reduction in host condition following infection with parasites from this location (Feis et al., 2016). The link between ROS production and ROS antagonist expression couples the expression patterns of parasites to those of the host and may be associated with to the phenotype of lowered body condition potentially resulting from oxidative stress. Further studies should therefore measure ROS production and oxidative stress directly to test this link on a functional level (Griendling et al., 2016).

Apart from the expression patterns of ROS pathways in the interaction between host and parasite, the differential expression of Chitotriosidase 1 homologs (Chit1) in mussels and cuticle proteins in M. intestinalis could be another example of a direct coupling of gene expression between hosts and parasites. As an essential component of the innate immunity against chitin-coated pathogens, Chit1 also regulates inflammatory processes (Elmonem, van den Heuvel, & Levchenko, 2016; Van Eijk et al., 2005). However, due to its digestive function in mussels (Birkbeck & McNerny, 1984; Lesser & Macmanes, 2016), oysters (Yang et al., 2015) and insectivorous scorpions (Fuzita et al., 2015), it is also another example of the dual roles of genes in digestion and immunity. Interestingly, the upregulation of Chit1 corresponded to the upregulation of a number of cuticle proteins in the parasites infecting Texel hosts (Figures 4, S1, and S2), suggesting that direct targeting of the parasite's chitin cuticle might have led to the parasite's response of strengthening the weakened exoskeleton with increased cuticle production. Since we only captured successful infections of parasites that were able to respond to the host attack, higher resistance in Texel mussels and higher infectivity of Texel parasites could be explained by successful expulsion of parasites that failed to counter the host chitinase immune effector by upregulation of their cuticle production. Hence, the coupling of host Chit1 and parasite cuticle proteins might characterize another direct interaction and help to explain how different evolutionary trajectories were realized along both invasion fronts (Feis et al., 2016).

5 | CONCLUSION

In this study, our dual-species transcriptomics approach elucidated some molecular underpinnings of the young co-evolutionary relationship between mussels M. edulis and their intestinal macroparasites M. intestinalis. The mechanistic interaction pathways described by several candidate genes partly matched the different
evolutionary trajectories of Mytilicola along the different invasion routes that have evolved over ~45 years since the introduction of the parasite, corresponding to roughly 45 host and probably around 100 parasite generations. Therefore, the observed changes occurred rapidly and had to rely on a sufficiently large standing genetic variation of both host and parasite populations or on more plastic mechanisms like gene regulation. Similar to many other nonmodel organism transcriptome studies (e.g., Gallardo-Escárate, Valenzuela-Muñoz, & Nuñez-Acuña, 2014; Patnaik et al., 2016; Qi et al., 2016; Yarra, Gharbi, Blaxter, Peck, & Clark, 2016), a significant proportion of assembled contigs could not be functionally annotated and we are aware that we may have missed novel functional aspects of co-evolutionary interactions. Our results based on homology annotation revealed that the direct interaction between host and parasites can generate specificity of general responses (ROS pathways) on the transcription level within few generations, illustrating the value of invasive species for the study of host-parasite co-evolution in nature. Although RNAseq is in principle a hypothesis-generating tool that cannot demonstrate causality, it can nevertheless be useful for identifying target genes and processes. Furthermore, transcriptional responses can help to characterize the observed phenotypes and describe co-evolutionary patterns. Future studies should focus on the functional significance of differential gene regulation in this system and try to separate the regulatory processes from adaptive evolution fixing underlying regulatory mutations. Here, we have mainly observed effects of M. intestinalis infestation and local adaptation on the effector side with our transcriptomics approach, but the underlying evolutionary changes, that is, the evolutionary changes at the DNA level that cause the patterns, are still elusive. Targeting the upstream steps of the involved pathways will therefore aid the search for key players in the co-evolutionary process in natural host-parasite systems.

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DATA ACCESSIBILITY

Raw RNAseq data are available in the BioProject PRJNA430138 on NCBI with BioSample Accession numbers SAMN08394988–SAMN08395065, which is linked to the Short Reads Archive study number SRP131446. Accession numbers SRR6513662-SRR6513728. Mussel M. edulis and parasite M. intestinalis assemblies and corresponding annotation tables are available upon request.

AUTHOR CONTRIBUTIONS

K.M.W. conceptualized the study. M.E.F. conducted the experimental work. M.E.F., U.J., A.L. and K.M.W. generated and analysed the data. M.E.F., U.J., A.L., P.C.L., K.M.W. wrote the manuscript.

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