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DOI:

[10.1111/mec.15886](https://doi.org/10.1111/mec.15886)

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Document Version

Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Zhang, C, Jones, M, Govaert, L, Viant, M, De Meester, L & Stoks, R 2021, 'Resurrecting the metabolome: rapid evolution magnifies the metabolomic plasticity to predation in a natural Daphnia population', *Molecular Ecology*, vol. 30, no. 10, pp. 2285-2297. <https://doi.org/10.1111/mec.15886>

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Resurrecting the metabolome: Rapid evolution magnifies the metabolomic plasticity to predation in a natural *Daphnia* population

Running title: Evolution magnifies metabolomic plasticity

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Abstract

Populations rely on already present plastic responses (ancestral plasticity) and evolution (including both evolution of mean trait values, constitutive evolution, and evolution of plasticity) to adapt to novel environmental conditions. Because of the lack of evidence from natural populations, controversy remains regarding the interplay between ancestral plasticity and rapid evolution in driving responses to new stressors. We addressed this topic at the level of the metabolome utilizing a resurrected natural population of the water flea *Daphnia magna* that underwent a human-caused increase followed by a reduction in predation pressure within ~16 years. Predation risk induced plastic changes in the metabolome which were mainly related to shifts in amino acid and sugar metabolism, suggesting predation risk affected protein and sugar utilization to increase energy supply. Both the constitutive and plastic components of the metabolic profiles showed rapid, likely adaptive evolution whereby ancestral plasticity and evolution contributed nearly equally to the total changes of the metabolomes. The subpopulation that experienced the strongest fish predation pressure and showed the strongest phenotypic response, also showed the strongest metabolomic response to fish kairomones, both in terms of the number of responsive metabolites and in the amplitude of the multivariate metabolomic reaction norm. More importantly, the metabolites with higher ancestral plasticity showed stronger evolution of plasticity when predation pressure increased, while this pattern reversed when predation pressure relaxed. Our results therefore highlight that the evolution in response to a novel pressure in a natural population magnified the metabolomic plasticity to this stressor.

Key words: adaptation; ancestral plasticity; crustaceans; metabolomics; rapid evolution of metabolic profiles; resurrection ecology

Introduction

Natural populations are facing rapid and strong environmental changes that ask for prompt responses to avoid local extinction. To realize these rapid responses to new environmental conditions, populations may rely not only on already present plastic responses (i.e. ancestral plasticity) but also on rapid evolutionary changes, whereby the latter can involve both evolution of mean trait values (i.e., constitutive evolution) and evolution of plasticity (Fox, Donelson, Schunter, Ravasi, & Gaitán-Espitia, 2019; Ghalambor, McKay, Carroll, & Reznick, 2007; Russell Lande, 2015). Against this background, important outstanding questions in evolutionary biology are what the relative contributions of the plastic and evolutionary responses are and how these relate to each other in determining rates of rapid adaptation (Fox et al., 2019; Ghalambor et al., 2015; Levis & Pfennig, 2016; López-Maury, Marguerat, & Bähler, 2008). There is still a debate going on regarding whether plasticity helps or hinders evolution. On the one hand, there is the view that ancestral plasticity may facilitate adaptive evolution, hence ancestral plasticity and the subsequent evolutionary response may covary positively (Espinosa-Soto, Martin, & Wagner, 2011; Scoville & Pfrender, 2010). On the other hand, the opposite has been hypothesized: that plasticity may slow down evolution by shielding traits from natural selection (Huey, Hertz, & Sinervo, 2003; Price, Qvarnström, & Irwin, 2003), resulting in negative covariation patterns between both. A major reason for why this controversy remains is the lack of compelling evidence from natural populations (Levis & Pfennig, 2016).

Because typically only the resulting phenotype after the joint action of plasticity and evolution can be studied, quantifying the ancestral plasticity is a key challenge when studying the interplay of plasticity and evolution in natural populations (Levis & Pfennig, 2016). Resurrection ecology offers a powerful solution when combined with common-garden

experiments where both the ancestral and derived genotypes of the same natural population that underwent a change in environmental conditions are reared in the absence and in the presence of the new selective agent. This combined retrospective approach allows reconstruction of microevolution of both trait means and their plasticity (Franks, Sim, & Weis, 2007; Weider, Jeyasingh, & Frisch, 2018). Resurrection ecology studies have been successfully applied to document evolutionary responses of natural populations to environmental stressors both at the levels of the phenotype (Hairston et al., 1999) and gene expression (Orsini, Spanier, & De Meester, 2012).

One way to advance insights in the interplay of ancestral plasticity and evolution in driving rapid trait adaptation is to focus on the underpinning molecular mechanisms (Fox et al., 2019). Such studies are very rare and limited to changes in gene expression (Ghalambor et al., 2015). Ignored so far in this context are responses in the metabolomes, the set of low-molecular-weight metabolites within organisms that is critical for unravelling the biochemical underpinnings of stress responses (Viant, Kurland, Jones, & Dunn, 2017). Metabolomic data can better reflect the activities at a functional level as metabolites are a downstream result of gene expression (Putri et al., 2013). Transcriptomic and proteomic data reveal the set of gene products being produced in the cell, data that represents one aspect of cellular function. Conversely, metabolic profiling can give an instantaneous snapshot of the physiology of that cell, and thus, metabolomics provides a direct "functional readout of the physiological state" of an organism (Hollywood, Brison, & Goodacre, 2006). Notably, metabolomics is also best suited to directly pick up any changes in free sugars which are expected under predation risk (meta-analysis of Rinehart & Hawlena, 2020), changes that other omics approaches (especially proteomics) would not be able to (directly) detect. Recently, metabolomics has been successfully applied to understand the plastic response to a variety of environmental

stressors (e.g., global warming, ocean acidification) (Calosi et al., 2017; Garreta-Lara, Campos, Barata, Lacorte, & Tauler, 2018; Mayor, Sommer, Cook, & Viant, 2015). We, however, lack information on the rapid evolution and its interplay with plasticity of metabolomic profiles in natural populations, which may provide unique insights in the interplay of ancestral plasticity and evolution at the molecular level.

Combining metabolomics with resurrection ecology and common-garden experiments, we tested the plastic and evolutionary responses of the metabolome to predation risk in a natural population of the water flea *Daphnia magna*. Predation is a potent selective pressure in natural populations. There is widespread evidence that predators may generate not only plastic but also rapid evolutionary responses in a wide array of phenotypic traits in prey populations including life history (Benard, 2004; Reznick, Shaw, Rodd, & Shaw, 1997), physiology (Hawlena & Schmitz, 2010), morphology (Eklöv & Svanbäck, 2017), and behaviour (Schoener, Losos, Kolbe, Lapiedra, & Leal, 2018). Yet, the metabolomic responses of prey (including both plastic and evolutionary responses) to predation remain unexplored. We capitalized on a unique *D. magna* population that experienced strong and opposing transitions in fish predation pressure through a period of ~16 years. First, there was a period with no fish predation (i.e. pre-fish subpopulation), followed by a period with high fish predation (i.e. high-fish subpopulation), and finally a period with relaxed fish predation (i.e. reduced-fish subpopulation) (Cousyn et al., 2001; Stoks, Govaert, Pauwels, Jansen, & De Meester, 2016). We have previously shown that the changes of fish predation pressure were associated with rapid, partly reversed evolution of phenotypic trait means and plasticity for a set of life history, morphology and behavioural traits (Stoks et al., 2016).

In the current study, we characterized the metabolomic profiles of 18 clones (6 clones from each of the three subpopulations differing in fish predation pressure) of *D. magna* both

in the absence and in the presence of fish kairomones. We first determined how predation risk changes the metabolomic profile of *D. magna*, i.e. which metabolites and metabolic pathways were affected by predation risk. We then partitioned the total changes of metabolomic profiles between successive periods in fish predation to quantify the relative contributions of ancestral plasticity, constitutive evolution, and evolution of plasticity (Stoks et al., 2016). Utilizing these data, we further tested whether and how plastic responses and evolutionary responses covaried with each other, testing the ideas whether metabolites that show the strongest ancestral plastic responses to fish kairomones also show the strongest evolutionary responses, or, alternatively, that these responses are negatively correlated (Huey, Hertz, & Sinervo, 2003). These analyses enabled us to get unique insights in the plasticity and evolution of the metabolome in a natural population in response to a novel stressor.

Materials and Methods

Resurrection of *Daphnia* clones and culture conditions

Daphnia magna clones were hatched from dormant eggs of three sediment layers of a shallow lake in Belgium (50°50' N, 4°39' E, Oud-Heverlee, Belgium). The three sediment layers match specific periods differing in fish predation pressure (Cousyn et al., 2001): the pre-fish period (1970 - 1972) with no fish present, the high-fish period (1976 - 1979) with high fish predation pressure because of intensive fish stocking (> 300 kg/ha), and the reduced-fish period (1988 - 1990) with relaxed fish predation pressure because of reduced fish stocking. An analysis of microsatellite variation indicated that the *Daphnia* from different periods can be considered as belonging to different subpopulations of one continuous population (Cousyn et al., 2001). For current study, six clones per period were used. To minimize interference from maternal effects, all clones were cultured under standard conditions for three generations prior to the experiment: up to 18 adults in 500 mL glass

vials, no fish kairomones, 20 °C, photoperiod 14:10 light:dark, daily fed *S. obliquus* (1.5×10^5 cells mL⁻¹, ~1.25 mg C L⁻¹) and with renewal of the culture medium every other day.

Experimental set-up

We tested for effects of fish kairomones, subpopulation and their interaction on the metabolomes in a full factorial experiment. The total design consisted of 6 clones \times 3 subpopulations \times 2 fish kairomone treatments \times 8 replicates = 288 experimental units. To manipulate fish predation risk, fresh medium containing fish kairomones was added daily. To prepare the medium, three fish (5-7 cm *Gasterosteus aculeatus* sticklebacks) were kept for 24 h in 20 L aerated and bio-filtered tap water. This fish-conditioned water was filtered twice (0.45 μ m) and diluted five times to obtain a final concentration of three fish per 100 L, which is known to generate strong responses in *D. magna* (Pauwels, Stoks, & De Meester, 2010; Zhang, Jansen, De Meester, & Stoks, 2016). The fish were fed *D. magna* daily in a separate bucket to avoid the presence of *Daphnia* alarm cues in the fish medium. The culture medium was refreshed every other day.

To obtain enough synchronized juveniles to start the experiment, for each clone we cultured ten to twelve *Daphnia* mothers from one grandmother. Cohorts of 16-18 juveniles of the pooled second brood of these mothers, all born within a 24 h interval, were used as experimental animals and cultured in 500 mL glass vials filled with 450 mL bio-filtered tap water. For each clone there were 8 replicates, hence 8 vials with 16-18 juveniles. To obtain relatively synchronised individuals for metabolomic profiling, one set of three individuals per vial were collected that had released their second clutch and had no visual signs of the third clutch in the brood pouch. Because the first clutch typically is small and not responsive to fish cues compared to the second clutch (Stoks et al., 2016), the physiological status and metabolome of the mother *Daphnia* might be more influenced by fish predation after release

the second clutch. When sampling, *Daphnia* were quickly rinsed with deionized water before transferring into a 1.5 ml centrifuge tube; the remaining water was gently removed with a glass Pasteur pipet. All samples were flash frozen in liquid nitrogen and then stored at -80 °C.

Metabolomic profiling

The metabolome of *Daphnia* samples was analysed at the Natural Environment Research Council (NERC) Biomolecular Analysis Facility at the University of Birmingham (UK). The metabolome of *Daphnia* samples were analysed using nano-electrospray ionization - direct infusion mass spectrometry (nESI-DIMS) as described by Southam *et al.* (2017) in their Supporting Information. Here, we applied nano-electrospray direct infusion mass spectrometry as this approach has been demonstrated to provide the required sensitivity when working with low biomass *Daphnia* samples and thereby allows running 3-4 technical replicates per sample (Southam et al., 2017; Taylor, Gavin, & Viant, 2018). Briefly, polar metabolites were first extracted from *D. magna* samples using a biphasic method. Then, all samples were analysed in both positive and negative ionisation modes using an Orbitrap Elite mass spectrometer (Thermo Fisher Scientific, Bremen, Germany) with a direct infusion, chip-based nano-electrospray ionization source (Triversa, Advion Biosciences, Ithaca, NY, USA). The data processing was done using the Galaxy online platform using the selected ion monitoring (SIM) stitching algorithm and data were acquired from mass to charge ratios (m/z) in the range 50–620 (Southam et al., 2017). After several steps (replicate filter, blank filter, sample filter, see details in Supporting Information), the processed data matrices were used for bioinformatics and statistical analyses.

Statistical analyses of metabolomic profiles

To assess the effects of exposure to fish kairomones and subpopulation on the metabolome of *D. magna*, we applied two parallel multivariate analyses: a principal component analysis

(PCA) coupled with two-way ANOVA and an ANOVA-simultaneous component analysis (ASCA). The results of both analyses were very similar and here we only show the PCA-ANOVA results (the ASCA results are in the Supplementary Information). PCA was conducted on the processed data matrices to assess the broad-scale variation between the two treatments using the PLS Toolbox (version 5.5.1, Eigenvector Research, Manson, WA, USA) within Matlab (version 7.8; The MathsWorks, Natick, MA, USA) following mean centring of the processed DIMS data. We extracted the first two PC axes for both ion modes; these explained 43.1 % (positive ion mode) and 42.9 % (negative ion mode) of the total variation. In order to test the effects of fish kairomones and subpopulation on the metabolome of *D. magna*, we then applied two-way ANOVAs on the generated PC scores in Statistica v12.0. In each analysis, the fish kairomone treatment, subpopulation and their interaction were included as fixed factors and clone was nested in subpopulation as a random factor. A significant effect of the fish kairomone treatment indicates plasticity, while a subpopulation effect indicates rapid evolution of the trait means, and a fish kairomone \times subpopulation interaction indicates rapid evolution of plasticity.

As we found strong fish kairomone \times subpopulation interactions on the metabolome, we then applied partial least squares discriminant analysis (PLS-DA) to each subpopulation separately to identify the specific metabolic responses to fish kairomones for each subpopulation. PLS-DA uses prior knowledge of the sample classes (here the fish kairomone treatments) to maximize separation of the metabolic profiles of the different classes and to derive predictive models (Nicholson, Connelly, Lindon, & Holmes, 2002). Internal cross-validation and permutation testing were employed to prevent over-fitting of the data (Westerhuis et al., 2008). Putative marker metabolites in response to fish kairomones for each subpopulation were screened using as criterion a Variable Importance in Projection (VIP)

threshold greater than 1 (Xuan et al., 2011). All putative marker metabolites for each subpopulation were compared to screen for the general and subpopulation-specific metabolite responses to fish kairomones. PLS-DA was conducted using in-house scripts with the PLS-Toolbox in Matlab.

In addition, changes in the intensities of individual m/z peaks were also assessed using t-tests for each subpopulation separately. All t-tests were corrected using a false discovery rate (FDR, Benjamini & Hochberg, 1995) of 5% to account for multiple testing and adjusted p-values are reported. Differences in the number of significantly changed peaks among subpopulations were tested using a chi-square test. The effects of fish kairomones and subpopulation on the metabolic rate was analysed using a two-way ANOVA in Statistica v12.0 (Stat-soft, Tulsa, OK, USA).

Metabolites annotation and pathway analyses

We used the MI-Pack software (Weber & Viant 2010) and the KEGG (Kyoto Encyclopedia of Genes and Genomes) platform to annotate the metabolites. The m/z peaks were assigned to possible, putative empirical formula(e) and KEGG compound names based on MI-Pack calculations. We then used MetaboAnalyst (Xia & Wishart, 2011) to analyse the metabolic pathways that were affected by fish kairomones. We put all putatively annotated KEGG compounds with VIP scores > 1 (based on the PLS-DA model including all three subpopulations) into MetaboAnalyst for metabolic pathway visualisation. Fisher's exact tests were used for over-representation analysis (Toyota, Gavin, Miyagawa, Viant, & Iguchi, 2016) and out-degree centrality was used for pathway topology analysis (Xia & Wishart, 2011). The FDR-corrected p values and impact values of all annotated pathways were plotted. Pathways were filtered based on the uncorrected p values ($-\log p > 0.5$) and impact value (> 0.2) as those pathways were considered as potentially affected (Ratnasekhar,

Sonane, Satish, & Mudiam, 2015). We also used “MS peaks to pathways” in MetaboAnalyst (Chong et al., 2018) to detect the affected metabolic pathways in each subpopulation separately.

Phenotypic trajectory analysis

We applied phenotypic trajectory analysis (PTA) to test whether the magnitude and direction of the multivariate plastic response of the metabolome to fish kairomones differed among subpopulations. This technique tests for pairwise differences between groups in multivariate plasticity (i.e. the phenotypic trajectories) by comparing the magnitude and the direction of the two-state multivariate reaction norms (Collyer, Adams, & Biology, 2007). PTA allows statistical testing for differences in magnitude and direction of phenotypic change by comparing observed values to distributions created from random pairs of trajectories obtained by permutations (Collyer et al., 2007). We compared the multivariate plasticity using all important metabolite peaks ($VIP > 1$) identified by the PLS-DA model including all three subpopulations.

The detailed methods of the PTA analyses are presented in the Supplementary Information. Briefly, we tested for differences in the magnitude and direction of the multivariate metabolomic change among the subpopulations using an extended R script of Adams & Collyer (Adams & Collyer, 2009) where the statistical model included fish kairomones, subpopulation and their interaction, and effects of clonal variation. To visualize the multivariate reaction norms, we conducted a principal component analysis, and plotted the scores on the first three, varimax normalized components. Note that these bivariate projection PCA plots cannot fully reflect the magnitudes and angles of the multivariate reaction norms as the PTA is conducted in a multi-dimensional trait space (Collyer et al., 2007).

The relative contribution of plasticity and evolution to metabolite changes

For both transitions between two successive periods differing in fish predation pressure we calculated the total peak change of important metabolites (VIP >1), i.e. the peak change as would be observed when comparing the *Daphnia* under the period-specific kairomone treatments (absence of fish kairomones in the pre-fish and reduced-fish periods, and presence of fish kairomones in the high-fish period), following the method described in (Stoks et al., 2016). We partitioned the total change for each important metabolite into three components: ancestral plasticity, constitutive evolution, and evolution of plasticity (see Figure 1 for details). Ancestral plasticity refers to the plasticity present in the older period of a given transition. Constitutive evolution refers to evolution of the mean in the ‘ancestral’ kairomone condition for a given transition between periods, hence in the absence of fish kairomones when going from the no-fish to the high-fish period and in the presence of fish kairomones when going from the high-fish to the reduced-fish period. Evolution of plasticity refers to the change in the slope of the reaction norm and is the remainder of the total trait change after subtracting ancestral plasticity and constitutive evolution from total phenotypic trait change (see Figure 1). Pairwise t-tests were conducted to compare the contributions of evolution of plasticity and of constitutive evolution to the total evolutionary change during both transitions.

The relationships between ancestral plasticity and evolutionary responses

For both transitions, we explored the relationships between ancestral plasticity and evolutionary changes (i.e., ancestral plasticity vs. total evolutionary changes, ancestral plasticity vs. evolution of plasticity, and ancestral plasticity vs. constitutive evolution). We therefore performed Pearson’s correlations between these components for all the identified important (VIP >1) metabolite peaks.

Results

General metabolomic responses

After mass spectral data processing, 2152 metabolites for the positive ion mode and 1519 metabolites for the negative ion mode were retained. The PCA results showed that fish kairomones induced a clear separation of the metabolomes along PC2 (Figure 2a, b). While there was also a separation along PC1, this was not linked to the treatments or subpopulations (Figure S1). Further two-way ANOVA on the PC2 scores showed significant effects of the fish kairomone and the fish kairomone \times subpopulation interaction (all $P < 0.001$, Table S1). These significant effects were confirmed by the ASCA model results (all $P < 0.0001$, Figure S2). Because of the significant fish kairomone \times subpopulation effects, PLS-DA analyses for each subpopulation were conducted separately, and the results confirmed that the fish kairomone treatment significantly changed the metabolomes in each subpopulation (all $P < 0.001$, Figure 2c-h, Table S2). Note that there was separation not only in LV1 but also in LV2 in the high-fish subpopulation (Figure 2e, f).

Metabolites annotation and pathway analyses

MI-Pack annotation showed 24.35 % (524 of 2152) of the peaks from the positive ion mode and 37.72 % (573 of 1519) of those from the negative ion mode could be assigned at least one molecular formula. Most of the annotated peaks were free amino acids and free sugars. The lists of the putatively annotated peaks are shown in the supplementary datasets 1 and 2. Analysing these putatively annotated peaks using MetaboAnalyst showed that predation risk generally affected four metabolic pathways (valine, leucine valine and isoleucine valine biosynthesis, D-glutamine and D-glutamate metabolism, purine metabolism, arginine and proline metabolism) for the positive ion mode and five pathways for the negative ion mode (starch and sucrose metabolism, biotin metabolism, galactose metabolism, ubiquinone and

other terpenoid-quinone biosynthesis, valine, leucine and isoleucine biosynthesis), which were mostly linked to amino acid and sugar metabolism (Figure 3).

When analysing each subpopulation separately, we found the most enriched pathways were different between subpopulations. For the positive ion mode, the pathway of ‘valine, leucine and isoleucine biosynthesis’ was mostly enriched in the pre-fish subpopulation while the pathway of ‘arginine and proline metabolism’ was mostly enriched for the high-fish and reduced-fish subpopulations (Table S3). For the negative ion mode, the pathway of ‘phenylalanine metabolism’ was mostly enriched for the pre-fish subpopulation, the pathway of ‘cysteine and methionine metabolism’ was mostly enriched for high-fish subpopulation, and the pathway of ‘alanine, aspartate and glutamate metabolism’ was mostly enriched for the reduced-fish subpopulation (Table S4).

Subpopulation-specific metabolomic profiles under predation risk

The metabolomic responses to fish kairomones differed strongly among subpopulations. Only a small part of the responsive peaks was shared by all three subpopulations: 15.9 % (176 of 1105, Figure 4a) for the positive ion mode, and 23.5 % (136 of 580, Figure 4b) for the negative ion mode. The total number of peaks whose levels were differentially modulated in response to fish kairomones differed among the subpopulations (positive ion mode: $\chi^2_2 = 278.39$, $P < 0.001$; negative ion mode: $\chi^2_2 = 79.40$, $P < 0.001$, Table S5). The high-fish subpopulation had the highest number of responsive peaks (Figure 4, Table S5), indicating that the metabolome of the high-fish subpopulation changed most strongly in response to fish kairomones.

Projecting the phenotypic trajectories onto the metabolomic PCA landscape showed that for both ion modes the magnitude of the multivariate metabolomic reaction norm was greater for the high-fish subpopulation than the pre-fish ($P < 0.001$ for both ion modes) and the

reduced-fish ($P = 0.026$ for the positive, $P < 0.001$ for the negative ion mode) subpopulations, while the latter two did not differ in magnitude (Figure 5a-d, Table S6). The direction of the multivariate plasticity differed considerably between the pre-fish subpopulation and the two other subpopulations (both $P < 0.001$). In contrast, the high-fish and reduced-fish subpopulations did not differ in the direction of multivariate plasticity for the positive ion mode ($P = 0.086$), and only differed slightly for the negative ion mode ($P = 0.047$) (Table S6).

Contributions of plasticity and evolution to total metabolomic changes in time

For the positive ion mode, ancestral plasticity had about an equal contribution to the total metabolomic change compared to the evolutionary components during both transitions in fish predation (47.3% in the first transition and 55.9% in the second transition, Figure 6a-b). Of the two evolutionary components, the contribution of evolution of plasticity was larger compared to constitutive evolution during both transitions (pre-fish to high-fish: 31.0% vs 21.7%, $t = -7.88$, $df = 517$, $P < 0.0001$; high-fish to reduced-fish: 24.6% vs 19.4%, $t = -5.41$, $df = 517$, $P < 0.0001$). The results were highly similar for the negative ion mode: ancestral plasticity had about an equal contribution compared with evolution during both transitions (46.5% and 48.8%). Of the two evolutionary components, the evolution of plasticity contributed more than constitutive evolution during pre-fish to high-fish transition (30.4% vs 23.1%, $t = -6.14$, $df = 405$, $P < 0.0001$) and high-fish to reduced-fish transition (30.0% vs 21.2%, $t = -7.46$, $df = 405$, $P < 0.0001$) (Figure 6c-d).

Relationships between ancestral plasticity and the evolutionary responses

For the positive ion mode, the ancestral plasticity in the pre-fish subpopulation correlated positively with the subsequent total evolution during the transition from pre-fish to high-fish ($R = 0.28$, $P < 0.0001$, Figure 7a). This pattern was driven by the evolution of plasticity ($R =$

0.43, $P < 0.0001$, Figure 7c), while the correlation of ancestral plasticity with constitutive evolution was negative ($R = -0.28$, $P < 0.0001$, Figure 7e). During the transition from high-fish to reduced-fish these patterns were reversed: the correlations with ancestral plasticity were negative for both total evolution ($R = -0.43$, $P < 0.0001$, Figure 7b) and evolution of plasticity ($R = -0.55$, $P < 0.0001$, Figure 7d), and positive with constitutive evolution ($R = 0.39$, $P < 0.0001$, Figure 7f). These patterns were largely similar for the negative ion mode (Figure S5).

Discussion

Using resurrection ecology, we provide unique evidence that changes in predation pressure can drive rapid evolution of metabolomes and their plasticity in a natural prey population. The high-fish subpopulation of *D. magna* evolved the strongest metabolomic response to predation risk thereby matching the changes in fish predation pressure across periods and the previously documented adaptive changes in life history, morphology and behaviour (Stoks et al., 2016). Key findings about the interplay of plasticity and evolution were (i) that ancestral plasticity and evolution contributed nearly equally in driving total metabolomic changes through time with the evolution of plasticity being the larger evolutionary component, (ii) and that the ancestral plasticity in the metabolome covaried positively with evolution of plasticity when predation pressure increased while this pattern reversed with subsequent relaxation of predation pressure.

Predator-induced plastic changes in metabolomic profiles

The mechanisms underlying the widespread trait responses to predation risk are still poorly understood (Mitchell, Bairos-Novak, & Ferrari, 2017). Using metabolomics, we found that the mostly affected metabolites of *D. magna* under predation risk were free amino acids and free sugars, and accordingly, amino acid metabolism (e.g., arginine and proline metabolism)

and sugar metabolism (e.g., starch and sucrose metabolism) pathways. These metabolic changes under predation risk may reflect either more breakdown or more synthesis of the associated polymers (proteins and carbohydrates). In the breakdown scenario, animals may face energy limitation under predation risk leading to an increased breakdown of the polymers to meet an increased energy demand. A recent meta-analysis of Rinehart & Hawlena (2020) suggested that predation risk indeed tended to decrease the carbohydrate and protein contents of prey to meet increased energy demands required to fuel stress responses. The here studied *D. magna* population reduced the RNA:DNA ratio when exposure to fish cues (Zhang et al., 2016), suggesting that the free amino acids changes may reflect an increased protein catabolism. Alternatively, in the synthesis scenario, *D. magna* may not be energy limited but re-allocate energy to other functions under predation risk. In support of this, *Daphnia* maintains feeding and assimilation rates (Beckerman et al., 2007; Stibor & Machacek, 1998) and even increases somatic growth rate (Stoks et al., 2016) under predation risk, and the experiment was performed under saturating food concentrations. The changed levels of free amino acids and sugars may therefore indicate more synthesis of proteins and carbohydrates. For example, protein synthesis (e.g. Hsp) is known to increase under predation risk to protect cells from protein malfunctioning in *D. magna* (Pauwels, Stoks, & De Meester, 2005; Pijanowska & Kloc, 2004). Further, exposure to fish kairomones has been shown to upregulate genes involved in protein folding (an important step for protein synthesis) in *D. magna* (Schwarzenberger, Courts, & von Elert, 2009).

Predation drove adaptive metabolomic evolution

By measuring the metabolomes of three subpopulations separated in time that belong to one continuous population and underwent strong changes in fish predation pressure, we could directly demonstrate rapid metabolomic evolution within a single natural population. The

three *D. magna* subpopulations did not seem to be equal in their ‘metabolic fingerprint’ not only in the presence (Figure S3) but also in the absence (Figure S4) of fish kairomones, suggesting rapid evolution of constitutive metabolic differences in this population. Besides constitutive evolution, we also observed rapid evolution of metabolomic plasticity to predation risk. Until now, it is poorly understood that whether and how the metabolome evolves in natural populations. As a notable exception, the marine snail *Littorina littorea* evolved different metabolomic responses to ocean acidification at much longer timescales associated with postglacial range expansion, which was linked to regional adaptation in physiology and life history (Calosi et al., 2017).

Two lines of evidence suggest that the rapid evolution of metabolomic responses to predation risk was adaptive. First, in line with the highest fish predation pressure being present in the high-fish subpopulation, this subpopulation showed a stronger metabolomic response to fish kairomones compared to the pre-fish and reduced-fish subpopulations. This was illustrated both by the high-fish subpopulation having the most metabolites responsive to fish kairomones (Figure 4) and the largest magnitude of the multivariate metabolomic reaction norm in response to fish kairomones (Figure 5). Notably, the fact that the high-fish subpopulation had the highest number of unique metabolites whose levels were significantly changed, might also have contributed to the separation in LV2 only being observed in the high-fish subpopulation (Figure 2e, f). That the predator-induced changes in the levels of these metabolites (particularly free amino acids and free sugars) are largest in the high-fish period, therefore suggests these *Daphnia* evolved to mobilize more energy to defend themselves against fish predation and associated predator-induced stress, which is consistent with the adaptive evolution of higher predator-induced plastic changes in their phenotype (life history, morphology and behaviour) (Stoks et al., 2016). Second, in line with fish

predators being only present in the high-fish and reduced-fish periods (Cousyn et al., 2001), these two subpopulations had a more similar direction of the multivariate metabolomic response under predation, while the pre-fish subpopulation differed to the high-fish and reduced-fish subpopulations. Notably, also the mostly affected metabolic pathway under simulated fish predation was the same in high-fish and reduced-fish subpopulations (arginine and proline metabolism), and differed from the one in the pre-fish subpopulation (valine, leucine and isoleucine biosynthesis, in the positive ion mode). As the free amino acids arginine and proline can be used by mitochondria as a metabolic fuel under stress (Raza et al., 2020), this may suggest that the high-fish and reduced-fish subpopulations evolved to better utilize energy when coping with predation risk. This pattern is also consistent with the multivariate reaction norms for life history, behaviour and morphology in a previous study (Stoks et al., 2016).

Noteworthy, the metabolomic responses to fish kairomones in the reduced-fish subpopulation did not fully convert back to those of the pre-fish subpopulation, similar to what was observed for life history traits (Stoks et al., 2016). This illustrates also at the metabolomic level that evolution in response to a new selective factor is not necessarily fully reversed when that selection factor is relaxed (Hairston et al., 1999; Lahti et al., 2009).

The interplay of plasticity and evolution in driving rapid metabolomic shifts through time

Partitioning the changes in metabolomic profiles during the two transitions in fish predation pressure showed that ancestral plasticity contributed approximately equal to the total changes in metabolite concentrations as evolution, i.e. the combination of constitutive evolution and evolution of plasticity (Figure 6). This is consistent with the important contribution of ancestral plasticity to the changes in life history and morphology in the studied system (Stoks

et al., 2016) and in other systems (Ghalambor et al., 2007). During both transitions, the evolution of plasticity was more important than constitutive evolution. This is in line with the expectation of evolutionary increases in plasticity in response to rapid increases of a novel selection pressure (Chevin & Lande, 2015; Lande, 2009; Robinson, 2013). It, however, deviates from earlier observations on phenotypic traits in this study system, for which evolution of plasticity played a more important role when fish predation increased whereas responses were more driven by constitutive evolution when predation was relaxed (Stoks et al., 2016). Thus, the current study indicated that the evolution of the metabolome and the evolution of phenotypic traits may partly be uncoupled. One possible reason for the discrepancy between the evolutionary patterns in the metabolome and the previously reported patterns in a set of 14 phenotypic traits, may be that the metabolome is thought to be linked to the phenome (the total sum of phenotypic traits) (Sardans, Peñuelas, & Rivas-Ubach, 2011), and we cannot exclude that other untested phenotypic traits actually follow the same pattern as the metabolome.

During the first transition, when the prey population experienced a strong increase in a novel selection agent (i.e. fish predation pressure), the ancestral plasticity and the evolution of plasticity were positively correlated. In other words, metabolites that showed a stronger plasticity to predation risk in the pre-fish subpopulation evolved to be even more plastic under increased fish predation pressure. This pattern corroborates the interpretation that the plastic response is adaptive and that evolution upon exposure to predation magnified the plastic responses of metabolomes to this stressor in this natural population. While the positive correlation between ancestral plasticity and evolution is consistent with the idea that ancestral plasticity may facilitate adaptive evolution (Fox et al., 2019; Ghalambor et al., 2007; Levis & Pfennig, 2016), an alternative explanation might be that plasticity in response to a stressor is

enhanced because it was adaptive and the stressor increased in strength. This latter interpretation is more in line with the pattern that we observe in the second transition.

Notably, when the selection was relaxed again during the second transition, the sign of the association reversed: the ancestral plasticity and evolution of plasticity were negatively correlated, indicating that metabolites with a stronger plasticity to predation risk in the high-fish subpopulation evolved to be less plastic under relaxed pressure. This suggests that the stronger metabolomic plasticity under predation risk as observed in the high fish period is costly and counter-selected against when predation pressure turns low. This matches the reduced plasticity for multiple life-history, morphology and behaviour traits in *D. magna* during the second transition (Stoks et al., 2016), as also has been observed in Trinidadian guppies when prey shift to a situation with relaxed predation pressure (Westrick, Broder, Reznick, Ghalambor, & Angeloni, 2019). Our results during the second transition match with the only other study reporting a correlation between ancestral plasticity and evolution at the molecular level. Trinidadian guppies transplanted from a site with high to low predation pressure also showed a negative association between ancestral plasticity (in this case of gene expression) and total evolution (Ghalambor et al., 2015). Ghalambor et al. argued the ancestral plasticity in the high-predation site to be maladaptive in the guppy system, and the most plastic transcripts to evolve reduced plasticity as a result of strong selection against non-adaptive plasticity (Ghalambor et al., 2015). Instead, in our study the positive association between ancestral plasticity and evolution of plasticity suggests adaptive plasticity in the high-predation period. Hence, our results indicate that in the *Daphnia* population the negative correlation between ancestral plasticity and evolution of plasticity upon relaxation of the predation pressure is in line with a true reversal of the response, with those metabolites that

evolved the strongest phenotypic plasticity during the first transition now showing the strongest reduction in plasticity.

The evolutionary responses in metabolite expression in our study population were driven by the evolution of plasticity. Constitutive evolution was a relatively minor component of the evolutionary response, and showed opposite covariation patterns with ancestral plasticity. Specifically, during the first transition metabolites that responded more plastically in the ancestral population showed less evolution in their mean levels in the absence of predation risk. This is suggestive of a pattern where ancestral plasticity can buffer evolution (Price et al., 2003), but in practice should be integrated with the evolution of plasticity. During the first transition, it thus seems that metabolites with higher ancestral plasticity show stronger evolution of plasticity but combine this with a change in main trait value in the ancestral environment that is opposite to the (change in) plasticity. Also, for constitutive evolution, the response pattern for the second transition reflects a reversal. Overall, our results show that the association between ancestral plasticity and evolution is complex, can depend on which component of evolution one studies, and might critically depend on whether one considers an increase in selection pressure or a release from this selection pressure. Our study also advocates for an integrated approach in which one goes beyond interpreting patterns from a one-directional study or across taxa and populations.

Conclusions

Our resurrection ecology study of evolution in a natural population provides unique input at the metabolomic level to the ongoing debate on the relationships between ancestral plasticity and subsequent evolutionary changes (Levis & Pfennig, 2016). We addressed two important outstanding questions (Fox et al., 2019; Ghalambor et al., 2015, 2007; Levis & Pfennig, 2016; López-Maury et al., 2008). First, we showed ancestral plasticity and evolution to

contribute nearly equally in driving total metabolomic changes through time. Second, we demonstrated that evolution of plasticity magnified the ancestral plasticity when a new selection pressure was imposed. Such insights are important to advance our ability to understand and predict how populations deal with the new and strong selection pressures which they are increasingly dealing with.

Acknowledgements

Financial support came from National Natural Science Foundation of China (42007229), China Postdoctoral Science Foundation (2019M662337), Post-doctoral Innovation Research Program of Shandong Province (236346), Research Grants from FWO Flanders (G.0943.15), and the KU Leuven Research Fund (C16/17/002). We thank Dr. Ralf Weber and Dr. Gavin Lloyd for helping with the metabolomic data analysis.

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Data accessibility

Metabolomics data have been deposited at dryad (doi.org/10.5061/dryad.vdncjsxtm).

Author contributions

701 CZ, LDM, MV and RS developed the concept of the study; CZ performed the experiment
702 with the input from MJ and MV for the metabolomics part; CZ analysed the data with the
703 input from MJ and LG. CZ wrote the first version of the manuscript; LDM and RS
704 contributed substantially to revisions and all authors agreed on the final manuscript.

Tables and Figures

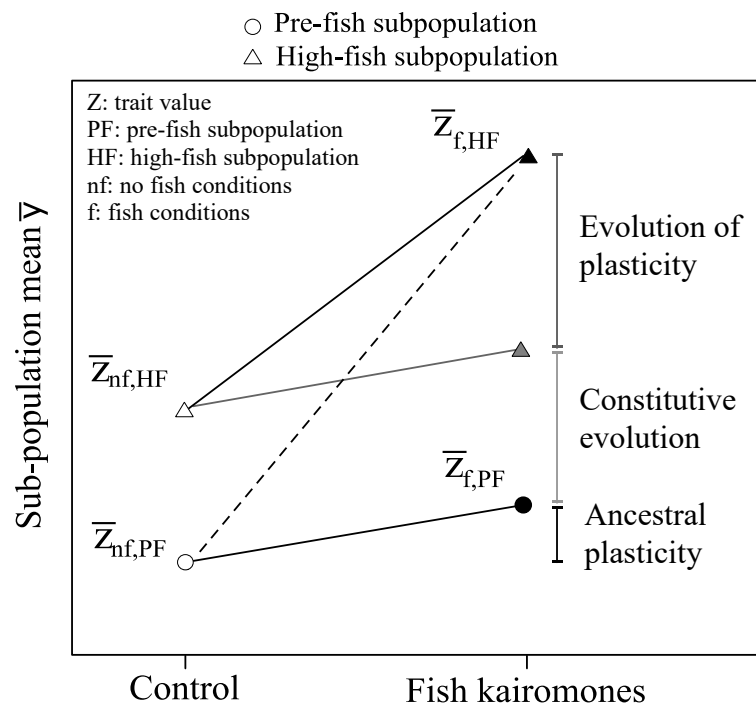


Figure 1 Methodology used to divide the total phenotypic trait change during a transition in fish predation pressure into three components: ancestral plasticity ($= \bar{Z}_{f,PF} - \bar{Z}_{nf,PF}$), constitutive evolution ($= \bar{Z}_{nf,HF} - \bar{Z}_{nf,PF}$) and evolution of plasticity ($= \bar{Z}_{f,HF} - \bar{Z}_{f,PF} - \bar{Z}_{nf,HF} + \bar{Z}_{nf,PF}$, see details in Stoks et al. 2016). Shown is the situation for the trait change between the pre-fish (●) and high-fish (▲) periods. White (black) symbols indicate subpopulation means in the absence (presence) of fish kairomones; the grey triangle represents the hypothetical trait value for the high-fish sub-population mean in the presence of fish kairomones in case only constitutive evolution and ancestral plasticity occur. The dashed line refers to the total trait change which is the trait change one would observe *in situ*.

Subpopulation	Control	Fish kairomones
Pre-fish	○	●
High-fish	△	▲
Reduced-fish	□	■

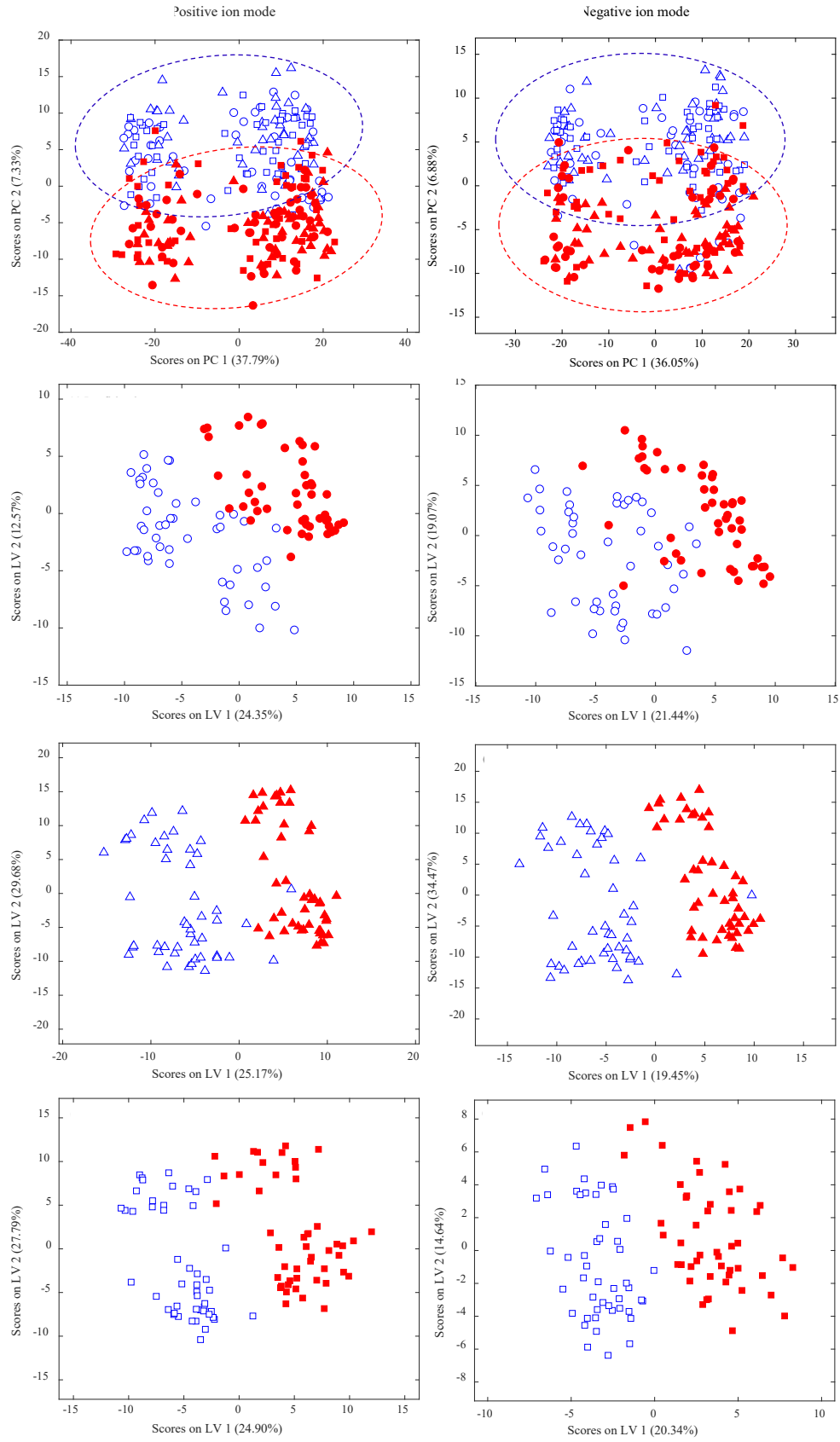


Figure 2 PCA and PLS-DA scores plots of positive (a, c, e, g) and negative (b, d, f, h) ion mode mass spectra describing the metabolomes of different subpopulations of *Daphnia magna* under control (blue colour) and fish predation risk (red colour). The ellipses of 95% confidence to the PCA plots were included in Figure 2a, b. LV1 and LV2 are the first two latent variables.

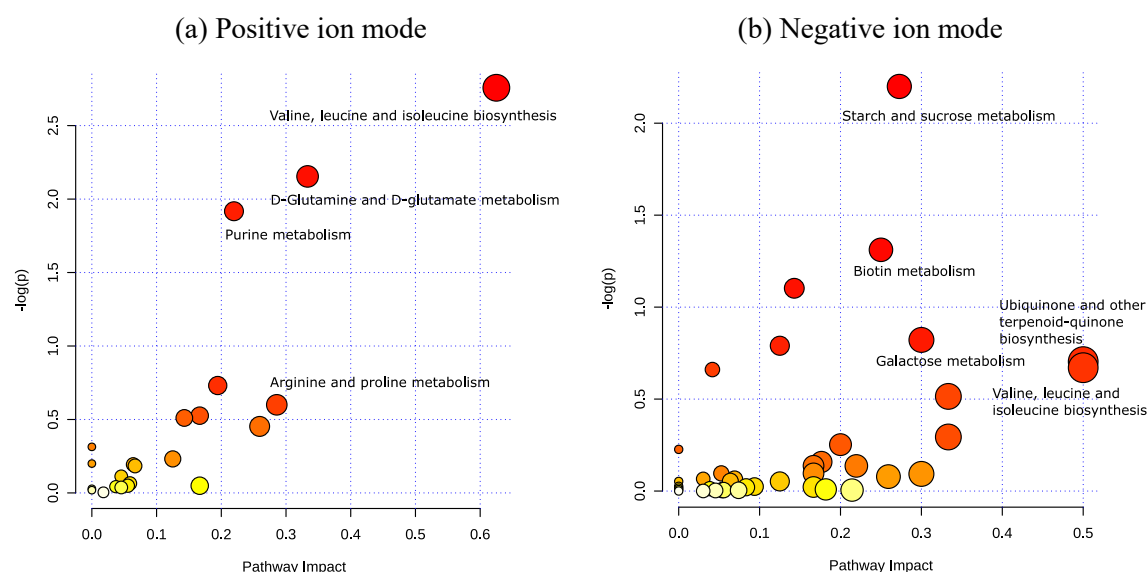


Figure 3 The identified metabolic pathways of *D. magna* that are most strongly reacting to fish kairomones. The circles represent pathways. The colour of each pathway is coded from white (lower impact value) to red (high impact value). The size of the circle is larger when pathway impact is higher. Only pathways with uncorrected p values ($-\log p > 0.5$) and impact value (> 0.2) were labelled in the figure as those pathways were considered potentially affected.

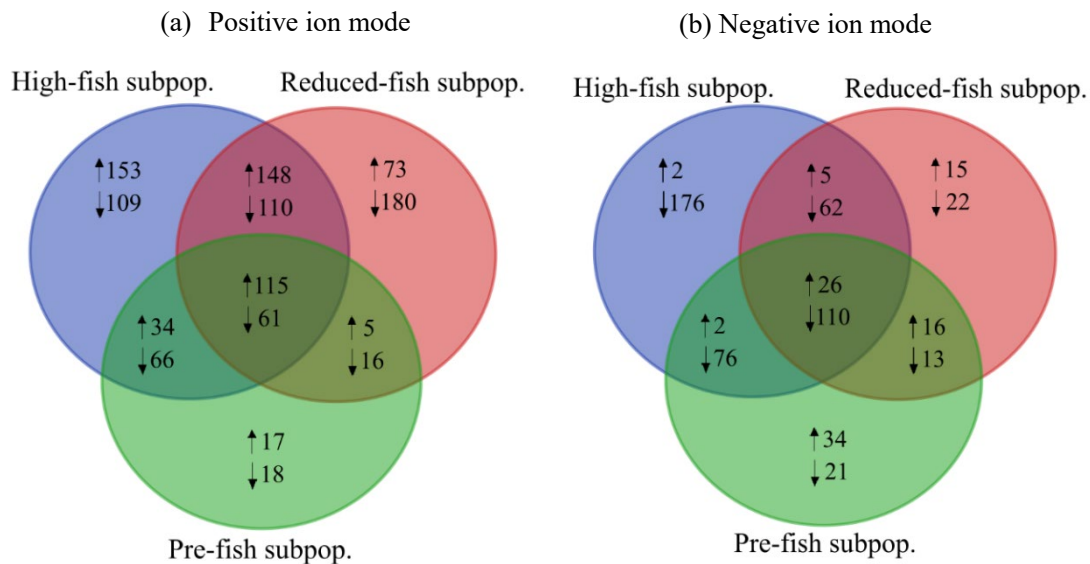


Figure 4 Venn diagram of the significantly up-modulated and down-modulated levels of metabolites in response to fish kairomones for each subpopulation of *Daphnia magna* for the (a) positive and (b) negative ion modes.

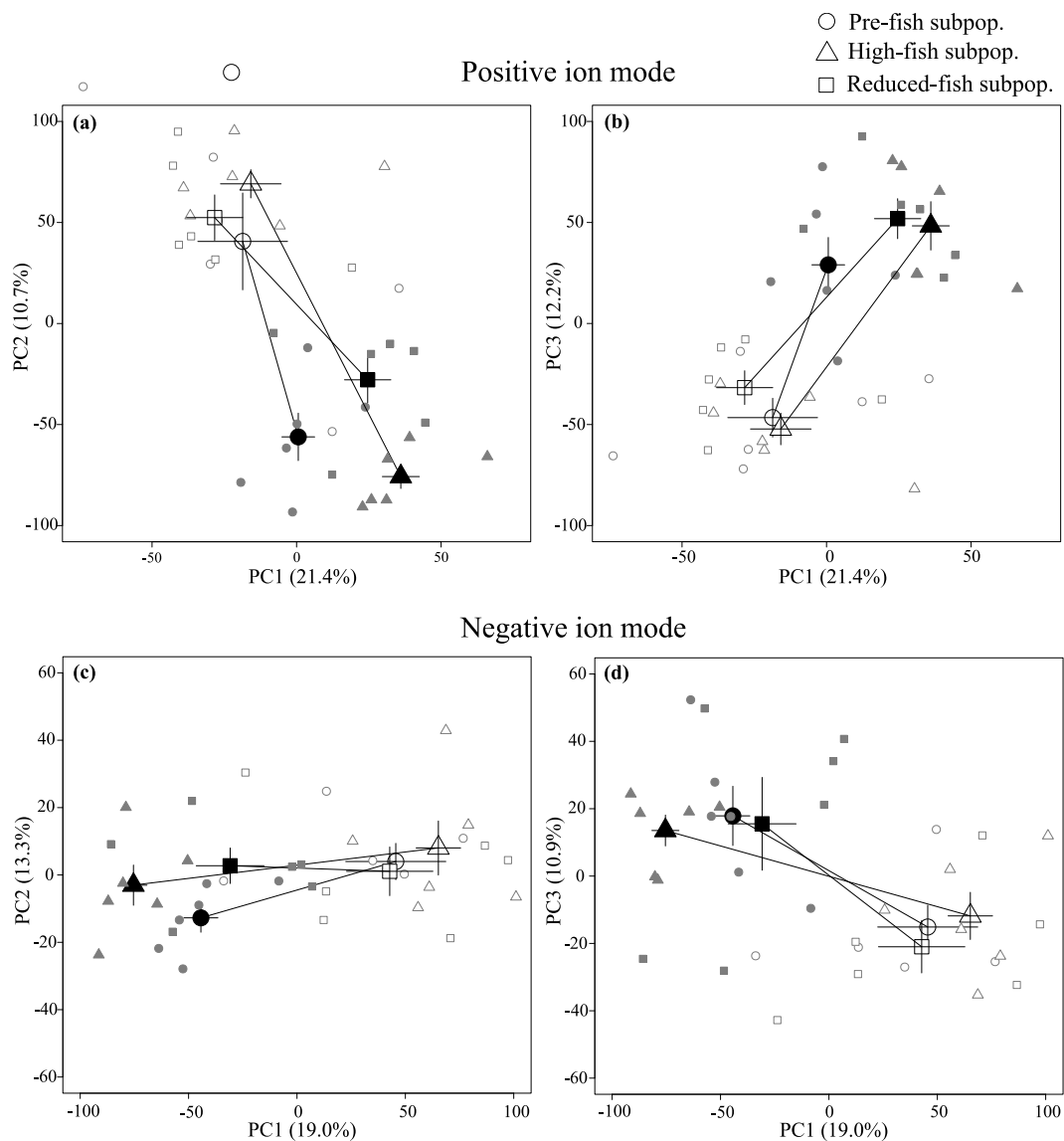


Figure 5 Multivariate metabolomic reaction norms representing the response to fish kairomones (open symbol: absence, filled symbol: presence) of the three subpopulations of *Daphnia magna*. Upper plots (a-b) show the patterns for the positive ion mode, lower plots (c-d) for the negative ion mode. Shown are patterns for PC1 and PC2 (a, c), and PC1 and PC3 (b, d).

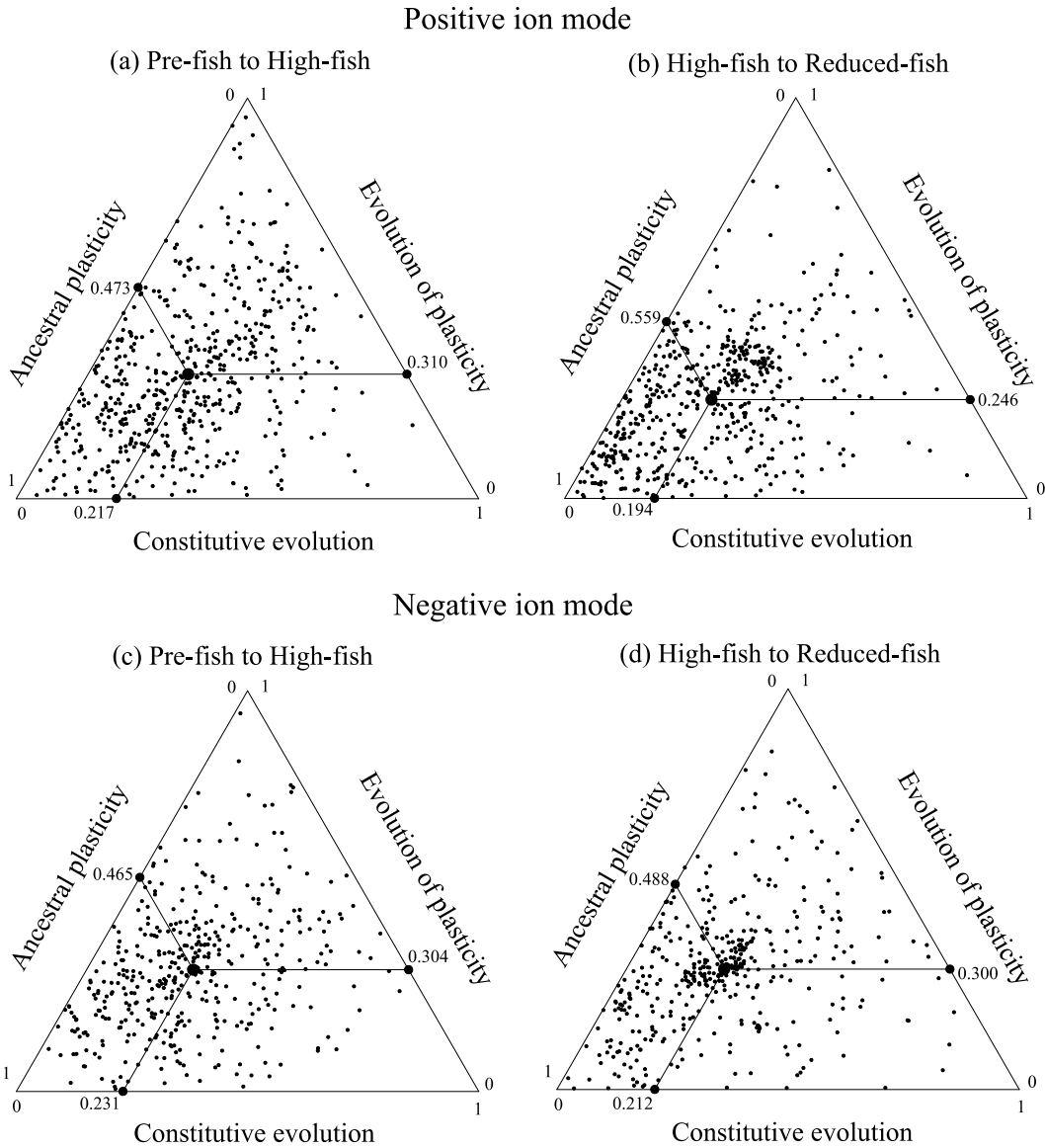


Figure 6 Relative contributions of ancestral plasticity, constitutive evolution and evolution of plasticity to the total changes in important metabolite peaks for (a, c) the transition from no fish to high fish predation and (b, d) the transition from high to reduced fish predation in the natural *D. magna* population. Partitioning was done separately for the positive ion mode (a, b; 518 peaks) and for the negative ion mode (c, d; 406 peaks). Shown are the results based on the partitioning method with contributions estimated using the additive method explained in Figure 1. Each small black dot represents a single metabolite peak. Large black dots represent the mean values of each component.

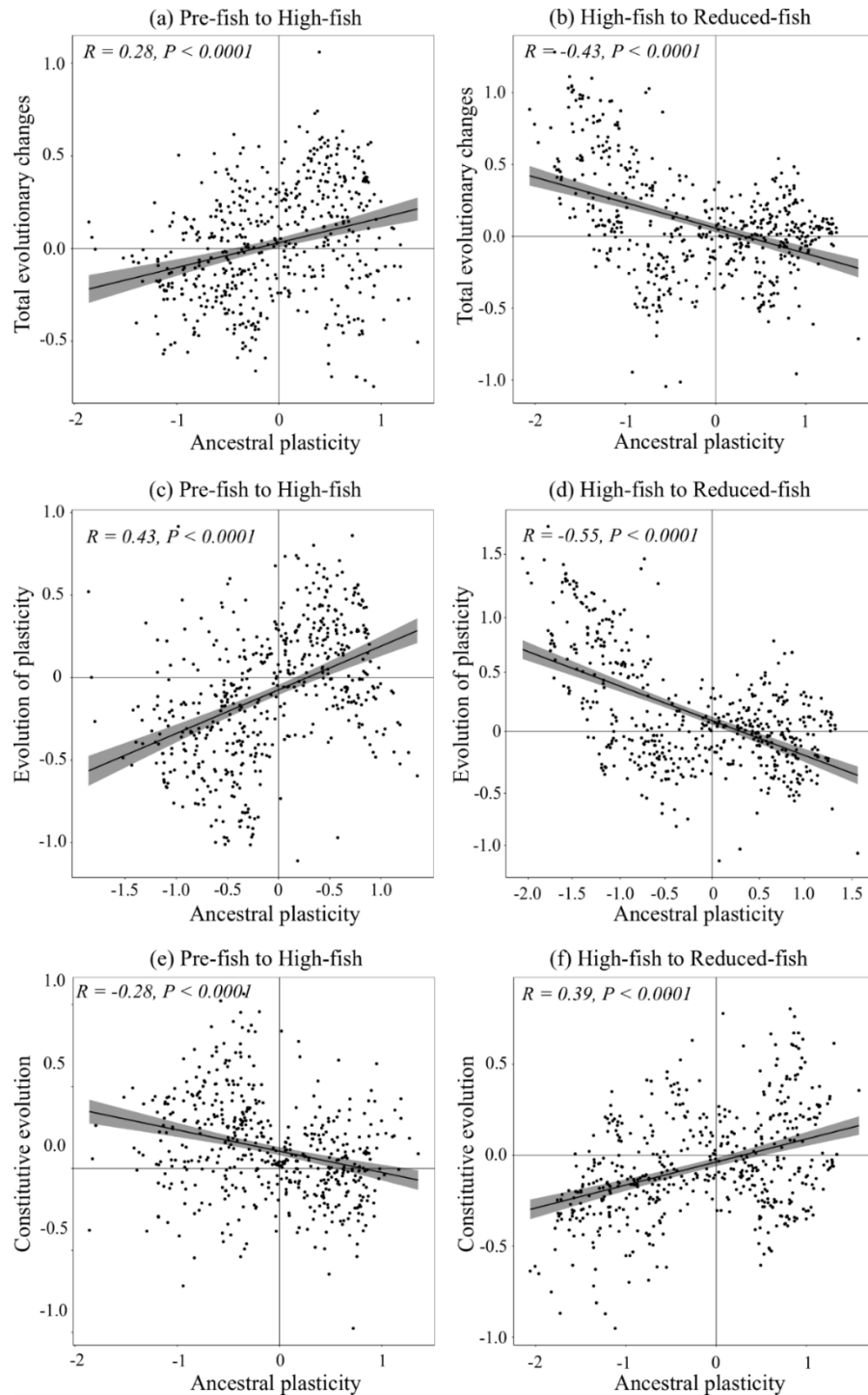


Figure 7 Relationships between ancestral plasticity and subsequent evolution of the metabolome: (a, b) total evolutionary changes; (c, d) evolution of plasticity; (e, f) constitutive evolution. Show are the patterns for both transitions in fish predation pressure for the positive ion mode (based on 518 peaks with $VIP > 1$). Each dot represents a single metabolite peak. Pearson's correlations with P -values are given.