Maternal dopamine exposure provides offspring starvation resistance in Daphnia

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Abstract

The neurotransmitter dopamine has been shown to play an important role in modulating behavioural, morphological and life-history responses to food abundance. However, costs of expressing high dopamine levels remain poorly studied and are essential for understanding the evolution of the dopamine system. Negative maternal effects on offspring size from enhanced maternal dopamine levels have previously been documented in Daphnia. Here, we tested whether this translates into fitness costs in terms of lower starvation resistance in offspring. We exposed Daphnia magna mothers to aqueous dopamine (2.3 mg/L or 0 mg/L for the control) at two food levels (ad libitum versus 30% ad libitum) and recorded a range of maternal life history traits. The longevity of their offspring was then quantified in the absence of food. In both control and dopamine treatments, mothers that experienced restricted food ration had lower somatic growth rates and higher age at maturation. Maternal food restriction also resulted in production of larger offspring that had a superior starvation resistance, compared to ad libitum groups. However, although dopamine exposed mothers produced smaller offspring than controls at restricted food ration, these smaller offspring survived longer under starvation. Hence, maternal dopamine exposure provided an improved offspring starvation resistance. We discuss the relative importance of proximate and ultimate causes for why D. magna may not evolve towards higher endogenous dopamine levels despite the fitness benefits this appears to have.

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The neurotransmitter dopamine has been shown to play an important role in modulating behavioural, morphological and life-history responses to food abundance. However, costs of expressing high dopamine levels remain poorly studied and are essential for understanding the evolution of the dopamine system. Negative maternal effects on offspring size from enhanced maternal dopamine levels have previously been documented in Daphnia. Here, we tested whether this translates into fitness costs in terms of lower starvation resistance in offspring. We exposed Daphnia magna mothers to aqueous dopamine (2.3 mg/L or 0 mg/L for the control) at two food levels (ad libitum versus 30% ad libitum) and recorded a range of maternal life history traits. The longevity of their offspring was then quantified in the absence of food. In both control
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Keywords
Phenotypic plasticity, life history, reaction norms, maternal effects, fitness

Introduction
Spatially and temporally heterogeneous environments promote phenotypic plasticity, the propensity of an organism to change its phenotype in response to changes in the environment (West-Eberhard, 2003). Under natural selection, adaptive phenotypic plasticity evolves such that the resulting reaction norm gives higher fitness across the changing environment (Lande, 2009). Phenotypic plasticity can also be transferred from mother to offspring such that the maternal response to the environment induces changes to the offspring reaction norm (Uller, 2008). Maternal effects can then impact offspring fitness and ultimately population dynamics (Benton et al., 2008). Hence, expression of adaptive reaction norms is essential to maintain high fitness as well as population viability in heterogeneous environments. This is especially the case for reaction norms that are expressed in response to food abundance, as changes in these can have strong effects on different components of life history (Boggs, 2009). In short-lived species for example, resource allocation to somatic maintenance (including survival) increases at the cost of growth and reproduction when food is limited (Lynch, 1989; Martínez-Jerónimo et al., 1994). Resulting changes in maternal resource allocation to offspring can ultimately influence offspring survival and reproduction (Enserink et al., 1995; Hafer et al., 2011; Saastamoinen et al., 2013).

At the physiological level, the neurotransmitter dopamine plays an important role as mediator of trait responses to food (see Barron et al., 2010 for a review on dopamine-mediated behavioural and morphological responses to food across taxa). Issa et al. (2020) showed that in addition to influencing morphological and behavioural traits, dopamine can regulate life-history responses to food abundance. Specifically, in the zooplankton species Daphnia magna, exposure to dopamine caused life-history reaction norms (age at maturation, fecundity) to change in a way that resulted in higher population growth rates (calculated from maternal life history traits) when food was limited (Issa et al., 2020). This happened without any apparent fitness costs at high food abundance. These observations raise the question of why endogenous dopamine levels do not evolve towards higher values. Issa et al. (2020) suggested this may be due to costs of high dopamine levels being paid by the offspring generation, which was not quantified in their study. Negative maternal effects on offspring size from dopamine treatments were detected. A smaller offspring size may have detrimental effects on offspring survival since body size in Daphnia is positively associated with filtering rates (Porter et al., 1983) and offspring survival under food limitation (Gliwicz & Guisande, 1992). Because offspring survival is a crucial component of maternal fitness, investigation of costs to offspring from enhanced maternal dopamine levels could give better insight into the selective forces shaping the evolution of the dopamine system.

In this study, we experimentally tested for effects of maternal dopamine exposure on offspring fitness. We exposed one generation (F₀) of D. magna to dopamine at high versus restricted food ration and starved the offspring (F₁) to measure their starvation resistance. Based on the previous findings of Issa et al. (2020), we predicted changes to the slopes of the life-history reaction norms under dopamine exposure that result in faster somatic growth and smaller offspring when food is limited. Moreover, we hypothesized that the smaller offspring originating from dopamine-exposed mothers would experience reduced survival under starvation.

Materials and methods
Study organisms

As previously outlined in Issa et al. (2020), ephippia containing resting eggs resulting from sexual reproduction of *D. magna* were collected in November 2014, in a pond at Værøy Island (1.0 ha, 67.687°N 12.672°E), northern Norway. Ephippial eggs were hatched in the laboratory and propagated clonally. For the current experiment, juveniles of a single clone (clone 49) of *D. magna* were asexually propagated for ten successive generations prior to use. A maximum of 30 individuals of *D. magna* were cultured in 2.5 L aquaria at 20 °C in a modified “Aachener Daphnien Medium” (ADaM) (Klütten et al., 1994, SeO2 concentration reduced by 50%), under long photoperiods (16h L: 8h D) using white fluorescent lamps. The medium was exchanged weekly and the animals were fed three times a week with Shellfish Diet 1800® (Reed mariculture Inc.) at a final concentration of $3.2 \times 10^5$ cells/mL.

Experimental design

For the F₀ generation, a similar experimental design as in Issa et al. (2020) was used, corresponding to a full factorial design with control, dopamine and two food rations (high versus restricted), with six 500 mL replicate beakers (non-aerated borosilicate beakers, Fisherbrand) for each of the four combinations (Fig. 1). Ten neonates (< 24 h old) were introduced into each beaker and kept at 20 °C under long photoperiods (16h L: 8h D) until maturation. The exposure concentration of dopamine (2.3 mg/L) was chosen for successfully inducing changes in *D. magna* growth (Issa et al., 2020; Weiss et al., 2015). An exposure protocol as previously outlined in Issa et al. (2020) was followed. Specifically, dopamine hydrochloride (Sigma-Aldrich, St. Louis, MO, USA) was first dissolved in 100 mL ultrapure water before dilution in ADaM to the desired exposure concentration. Controls containing only ADaM medium were performed parallel to the exposure replicates. The medium was renewed in all replicates (N = 24) three times a week, and the animals were fed at each renewal event with Shellfish Diet 1800® at a final concentration of $2.88 \times 10^5$ cells/mL (ad lib at 20 °C) for the high food ration and $8.6 \times 10^4$ cells/mL (30% ad lib at 20 °C) for the restricted food ration.

All replicates were checked daily for mature individuals (defined as having their first clutch of eggs released into the brood chamber). Mature individuals were removed from the beakers, photographed for size measurements (see section 2.3 below) and placed individually in 2 mL non-aerated wells containing ADaM at 10 °C to give birth. Seventeen of the 240 individuals in the beakers were found to be males and were thus not transferred (Fig. 1). To prevent resulting neonates from obtaining food upon birth, no food was added to the wells. The wells were checked daily and once neonates had been released (i.e. < 24 h old, F₁ generation), two individuals were randomly sampled from each well, photographed for size measurements (see section 2.3 below) and transferred individually to new 2 mL non-aerated wells containing ADaM (no food) at 10 °C. The medium was not exchanged, and individuals were checked daily to record mortality. The low exposure temperature was chosen to slow down metabolism and increase longevity under starvation, thus making it more likely to accurately quantify any variation in offspring longevity under the chosen frequency of observations.

Sampling procedure and measurements of life-history traits

Immediately prior to dopamine exposure on day 0 of the experiment, a random sample of thirty F₀ neonates were photographed for body length measurements (mm, measured from the upper margin of the eye to the junction of the carapace and spine, using ImageJ v1.52a, National Institutes of Health, Bethesda, MD). Individual measurements of body length at first reproduction for the F₀ generation (after releasing their first clutch of eggs) and at birth for the F₁ generation was measured in the same way. All body length measurements (mm) were converted to dry mass (mg) using the equation by Yashchenko et al., (2016): $\text{Dry mass} = 0.00535 \times \text{Body length}^{2.72}$. The somatic growth rate in the F₀ generation was calculated for each individual as: $\text{Somatic growth rate} = \frac{\ln(\text{dry mass}_\text{end}) - \ln(\text{dry mass}_\text{start})}{\text{duration}}$, where dry mass start is the average dry mass (in mg) at the neonatal stage (day 0), dry mass end is the individual dry mass (in mg) at maturation, and duration is the number of days between the two stages.

As in Issa et al. (2020), conductivity (WTW LF 330 conductivity meter), pH (WTW pH 340i) and dissolved oxygen (WTW Multi 3410 multiprobe meter) were measured. This was done on two separate occasions.
throughout the experiment (Table 1), after medium renewal, in the new exposure solutions and ADaM medium used for the controls (N = 4; 2 samples collected in total from each of the dopamine and control treatments). Simultaneously, the new exposure solutions and ADaM medium were sampled for dopamine analysis (N = 4; 2 samples collected in total from each of the dopamine and control treatments). The samples were stored at -80 °C for a maximum of five months after collection (during closure of the labs due to the Covid-19 outbreak), prior to dopamine analysis. Two complementary sample preparation protocols were used: (1) dilute-and-shoot; and (2) liquid-liquid extraction. Subsequent analysis was performed by ultra-performance liquid chromatography coupled to a triple quadrupole mass analyser (UPLC-MS/MS). This analysis was done as described in Issa et al. (2020), where single reaction monitoring (SRM) chromatograms were monitored for dopamine. Typical SRM chromatograms for dopamine in the analysed samples are depicted in Appendix 1. Over the course of the experiment, pH, conductivity and dissolved oxygen were within the recommended range for testing of chemicals in *D. magna*, according to OECD guidelines (OECD, 2012). The conductivity remained at 1.6 mS/cm, mean dissolved oxygen at 8.9 mg/L and pH at 8.0 across treatments, whereas measured average concentrations of dopamine were ca. 2% of their nominal concentrations (Table 1). Much lower than expected concentrations of this compound were likely caused by degradation during unforeseen long-term storage due to the Covid-19 outbreak.

**Statistical analyses**

All statistical analyses and graphic illustrations were performed in R v. 3.6.3 (R Core Team, 2020). We tested whether the slopes of the reaction norms of the measured life-history traits in response to food abundance differed among dopamine treatments (control versus dopamine exposure). To assess this for dry mass at maturation and somatic growth rate, we used linear mixed effects models (LME, implemented using the `lme` functions in the package `nlme`, Pinheiro et al., 2020), with fixed effects of treatment and food ration and the interaction between these, and with beaker id as a random effect. For offspring dry mass, LME models were fitted with treatment, food ration and their interaction as fixed predictor variables, and maternal id nested in beaker as a random predictor variable. Finally, we tested the effects of treatment, food ration and their interaction on age at maturation and offspring longevity, using Poisson generalized linear mixed-effects (GLMM, implemented using the `glmer` function in the package `lme4`, Bates et al., 2015). For these latter analyses, beaker id and maternal id nested in beaker id were included as random effects, respectively. For LME models, the VarIdent command from the `nlme` package was used to allow residual variance to differ among treatments, food ration and the two-way interaction between these (Pinheiro & Bates, 2000). The appropriate random structure with respect to this was chosen based on AICc comparisons. GLMM models were tested for overdispersion and their Pearson and deviance residuals were checked for patterns and lack of fit. For all analyses, fits of alternative models with different fixed effects structures were compared using AICc.

**Results**

All life-history traits were influenced by both food ration and dopamine treatments (Table 2). Most of them responded in the same direction to a change in food ration for both control and dopamine treatments. Specifically, somatic growth rate increased with higher food ration, whereas age at maturation, offspring dry mass and offspring longevity decreased (Fig. 2, Table 3). For dry mass at maturation there was a strong interaction between food ration and dopamine treatment (Table 2). While dry mass at maturation decreased with increasing food ration in the control treatment, the opposite pattern was observed in the dopamine treatment (Fig. 2C, Table 3). The steepness of the food ration reaction norms depended on dopamine treatment for somatic growth rate and offspring dry mass (Table 2). Specifically, dopamine exposure increased somatic growth rate, but this was more pronounced at high food ration (Table 3), causing the reaction norm to be steeper for the dopamine exposed group compared to the control (Fig. 2A). For offspring dry mass, treatment had no effect at high food ration, whereas exposure to dopamine reduced offspring dry mass at restricted food ration (Table 3). Thus, for this trait, the reaction norm was steepest for the control group (Fig. 2D). In contrast, exposure
to dopamine lowered age at maturation to an equal extent across food rations (Fig. 2B, Table 2). Similarly, the effect of dopamine treatment on offspring longevity did not depend on the maternal food ration (Table 2). Rather, exposure to dopamine increased offspring longevity across food rations (Fig. 2E).

Discussion

In a previous study, Issa et al. (2020) investigated the role of dopamine in shaping life history responses to food abundance in *D. magna*. This was done both through aqueous exposure to dopamine and to the antidepressant bupropion, a dopamine reuptake inhibitor. Both treatments led to higher population growth rates (calculated from maternal life history traits) when food was restricted, without any apparent costs to fitness. The higher dopamine exposure resulted however in smaller offspring, which could potentially perform worse particularly when facing restrictions in food availability (Gliwicz, 1990). Since offspring survival is a crucial component of maternal fitness, this may represent a fitness cost of higher maternal dopamine levels.

In the current study we tested this by exposing F₀ *D. magna* to aqueous dopamine and quantifying life-history responses of mothers (F₀) to food abundance as well as the starvation resistance of their offspring (F₁).

Similar to the findings of Issa et al. (2020), offspring size was larger in mothers that had experienced restricted food abundance, a pattern commonly observed in *Daphnia* (Garbutt & Little, 2014; Glazier, 1992), as well as in other organisms (Reznick et al., 1996; Vijendravarma et al., 2010). This response was in the same direction for both dopamine and control treatments but was considerably steeper for the control treatment. The latter arose as a consequence of dopamine exposure causing a reduction in offspring size when mothers experienced restricted food ration, but not when mothers received *ad lib* food. In general, a larger investment in offspring size when food is limited is expected to boost offspring survival and fitness (Gorbi et al., 2011; Tessier & Consolatti, 1989). In support of this, we found that the larger offspring of mothers having experienced restricted food ration survived better under starvation across treatments. Mothers that experience low food abundance tend to produce offspring with a larger maternal lipid reserve, which can increase offspring starvation resistance (Tessier et al., 1983). In contrast to our hypothesis, however, maternal exposure to dopamine did not come at a cost to offspring longevity. Surprisingly, offspring in the dopamine treatment survived longer than controls across food rations. This was true even for the offspring from mothers that had experienced restricted food rations, where offspring size was smaller in the dopamine treatment than in the controls. Hence, enhanced maternal dopamine levels increased rather than decreased offspring survival.

Unlike the effects of dopamine exposure on offspring longevity, its effects on maternal life-history reaction norms were overall as predicted based on the findings of Issa et al. (2020). Specifically, at restricted food ration, somatic growth rate decreased, and maturation was delayed. However, dopamine exposure resulted in faster growth and earlier age at maturation across food rations compared to the control. Higher dopamine levels have been shown to promote cell proliferation and/or increase cell volume, explaining the positive effect of dopamine exposure on somatic growth rate (Huet & Franquinet, 1981; Weiss et al., 2015). As in Issa et al. (2020), a smaller adult size was observed at restricted food ration in the dopamine treatment, with potential costs to adult survival and reproduction (Cleuvers et al., 1997; Lampert, 2001), as body size in *Daphnia* positively correlates with the ability to satisfy metabolic requirements at low food levels. Contrastingly, at high food ration, dopamine exposure increased adult size, and hence *Daphnia* competitiveness (Brooks & Dodson, 1965), compared to the control. The significant effects of dopamine treatment on adult size and somatic growth rate (and consequently age at maturation) at high food ration were surprising, given that treatment effects were expected to occur at restricted food ration only (Issa et al., 2020). There were however some differences in the setup between the current study and the study by Issa et al. (2020) that may potentially explain this: 1) in the present study, we performed exposure in groups, thus allowing for interactions between individuals, whereas Issa et al. (2020) exposed animals individually; 2) the present study used a different clone than Issa et al. (2020). Previous research shows that antidepressant effects on life history traits of *Daphnia* species can vary between clones, if they differ in their growth and reproductive performance (Campos et al., 2012). Yet, although the exact patterns may depend on clonal identity and experimental protocols, the results of the present study support the overall conclusion of Issa et al. (2020)
that important life-history responses to food abundance are shaped by the dopamine system.

The observed increase in *Daphnia* growth and advanced timing of reproduction from elevated dopamine levels may also be interpreted as a stress response. Indeed, evidence shows that fast species (i.e. short-lived species, producing many offspring early in life), such as *Daphnia*, respond to stressful environments by accelerating their life cycle (Rochet et al., 2000; Trippel, 1995). Higher dopamine levels may stress individuals through dopamine oxidation. Elevated intracellular dopamine levels can promote the intracellular oxidation of dopamine, by increasing the production of reactive quinones and free radicals for a given rate of oxidation (Miyazaki & Asanuma, 2008; Sun et al., 2018). In addition, increased extracellular dopamine can overload the antioxidant capacity in the extracellular space, causing dopamine oxidation (Blesa et al., 2015; LaVoie & Hastings, 1999; Miyazaki & Asanuma, 2008). In the case of limited investment in antioxidant defence, dopamine oxidation can then lead to oxidative stress, which can potentially reduce longevity (Ishii et al., 1998; Moskovitz et al., 2001). On the other hand, increased investment in antioxidant defence may lower investment in immune defence (Takahashi et al., 2017), ultimately increasing the susceptibility of individuals to diseases and parasites. A higher investment in antioxidant defence may also lower investment in reproduction (Speakman & Garratt, 2013). There may also be ecological costs of rapid growth associated with enhanced dopamine levels, due to biotic interactions. One such cost may be higher predation risk from faster growth (Urban, 2007), as individuals become more exposed to predators from increased feeding (Lankford et al. 2001; Stoks et al., 2005). Thus, whether through oxidative stress, or increased investment in antioxidant defence, dopamine oxidation may pose a proximate restriction for why *Daphnia* do not produce more endogenous dopamine. This in addition to more ultimate restrictions in the form of reduced survival, from oxidative stress, reduced immunity or increased predation risk, as well as lowered reproduction, may be important to explore in order to understand the evolution of the dopamine system.

In summary, maternal dopamine exposure boosted *Daphnia* offspring survival in addition to accelerating its life cycle. Altogether, these results emphasize the important role of dopamine as a regulator of life-history responses to food abundance but leave open the question of why *D. magna* do not evolve towards higher endogenous dopamine levels despite the apparent fitness benefits. A potential proximate cause may be that higher dopamine levels promote dopamine oxidation. More ultimate causes involve the potential costs of dopamine oxidation to reproduction and survival, as well as ecological costs of rapid growth due to biotic interactions (predation). Hence, further understanding of the evolution of the dopamine signalling system may require a combined investigation of ultimate and proximate causes.

**Author contributions**

SI and SE conceived the study. SI, SE and VLBJ designed the methodology. SI, SC and AGA collected the data. SI analysed the data. SI led the writing of the manuscript with contributions and comments from all authors. All authors gave final approval for publication.

**Data availability statement**

All data used for this work are available on figshare, https://figshare.com/articles/dataset/Dataset_for_-dopamine_manipulated_daphnia/18030899

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Table 1. Water quality measurements from study on life-history responses to food ration and dopamine exposure in *Daphnia magna*.

| Sampling event | Treatment | Dopamine (mg/L) | Conductivity (mS/cm) | Dissolved oxygen (mg/L) | pH |
|----------------|-----------|-----------------|----------------------|------------------------|----|
| 1              | Dopamine  | 0.052           | 1.64                 | 8.83                   | 7.95|
|                | Control   | 0               | 1.653                | 8.93                   | 8.02|
| 2              | Dopamine  | 0.056           | 1.527                | 8.97                   | 7.98|
|                | Control   | 0               | 1.541                | 9.02                   | 8.01|

Table 2. Model comparison of candidate models for testing effects of treatment (control versus dopamine) and food ration (high versus restricted) on somatic growth rate, age at maturation, dry mass at maturation, offspring dry mass and offspring longevity in *D. magna*. The least complex model within 2 ΔAICc is given in bold. vI refers to the varIdent function.

| Response variable | Model | Model |
|-------------------|-------|-------|
| Somatic growth rate | Somatic growth rate | Somatic growth rate |
| Fixed             | **Food: Treatment** | **Food: Treatment** |
|                   | Food + Treatment | Food + Treatment |
|                   | Food            | Food            |
|                   | Treatment       | Treatment       |
Table 3. Parameter estimates from the best models (Table 2) describing the response of life-history traits to food ration and dopamine exposure in *Daphnia magna*.

| Response variable | Final model | Parameter | Estimate ± SE |
|-------------------|-------------|-----------|---------------|
| Somatic growth    | Food: Treatment + (1|Beaker) | Intercept | 0.24 ± 0.006 |
| rate              | vI (Treatment) + (1|Beaker) | Dopamine treatment | 0.02 ± 0.01 |
|                   | vI (Food: Treatment) + (1|Beaker) | High food | 0.09 ± 0.009 |
|                   | vI (Food) + (1|Beaker) | High food: Dopamine treatment | 0.03 ± 0.01 |
| Age at maturation (days) | Food + Treatment + (1|Beaker) | Intercept | 2.38 ± 0.04 |
|                   | Food: Treatment + (1|Beaker) | Dopamine treatment | -0.12 ± 0.05 |
|                   | vI (Treatment) + (1|Beaker) | High food | -0.37 ± 0.05 |
| Response variable | Final model | Parameter | Estimate ± SE |
|-------------------|-------------|-----------|---------------|
| Dry mass at maturation (mg) | Food: Treatment + (1|Beaker) | Intercept | 0.07 ± 0.001 |
| | | Dopamine treatment | -0.007 ± 0.002 |
| | | High food | -0.006 ± 0.002 |
| | | High food: Dopamine treatment | 0.01 ± 0.003 |
| Offspring dry mass (mg) | Food: Treatment + vI (Treatment) + (1|Beaker/Maternal) | Intercept | 0.003 ± 0.0001 |
| | | Dopamine treatment | -0.0004 ± 0.0001 |
| | | High food | -0.0007 ± 0.0001 |
| | | High food: Dopamine treatment | 0.0004 ± 0.0002 |
| Offspring longevity (days) | Food + Treatment + (1|Beaker/Maternal) | Intercept | 1.67 ± 0.04 |
| | | Dopamine treatment | 0.17 ± 0.05 |
| | | High food | -0.12 ± 0.05 |

**Figure captions**

**Fig. 1.** Schematic diagram of the experimental design. The F₀ generation was kept at restricted (R) and high (H) food ration for both control and dopamine treatments (2 x 2 design) from birth until maturity, with six replicate beakers per treatment combination and with each replicate initially containing ten individuals. Due to the occurrence of males in the F₀ generation, total sample sizes, given as N, varied among treatments. Individuals in the F₁ generation were kept without food in separate wells and were not exposed to dopamine.

**Fig. 2.** Effect of food ration on F₀ and F₁ traits in *D. magna* in the dopamine and control treatments. (A) Somatic growth rate, (B) age at maturation (days), (C) dry mass at maturation (mg), (D) Offspring dry mass (mg) and (E) offspring longevity (days). Error bars give 1SE.
Appendix 1

Dopamine was confirmed with the highest degree of certainty since 4 SRM transitions were monitored instead of a single transition (Fig. A1). Each SRM transition denotes a characteristic fragmentation pathway of the target molecule. We considered a positive finding in the samples when all four SRMs demonstrated a positive signal in the expected retention time for dopamine. The control samples did not present any signal in any of the SRM transitions, denoting non-detectable concentrations.

Fig. A1. Single Reaction Monitoring (SRM) chromatograms for dopamine in the four samples analysed; transitions monitored were: 154.1 > 137.0, 154.1 > 90.9, 154.1 > 119.0 and 154.1 > 65.0 m/z. Sampling dates were 1 April 2020 (denoted by 1.4) and 6 April 2020 (denoted by 6.4)