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Dietary antioxidants, food deprivation and growth affect differently oxidative status of blood and brain in juvenile European seabass (*Dicentrarchus labrax*)

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\textbf{Running title:} Early life and oxidative stress
Compensatory growth may increase molecular oxidative damage, which may be mitigated through the intake of dietary antioxidants. However, dietary antioxidants may also reduce concentration of antioxidant enzymes, which have a key role in regulating the oxidative status. Here we investigated whether feeding on a diet rich in antioxidants (vitamin E) enables juvenile European seabass (Dicentrarchus labrax) to catch up after a period of food deprivation with negligible effects on the oxidative stress to blood and brain as compared to fish feeding on a normal diet (i.e., not enriched in antioxidants). The results show that a higher intake in antioxidants favoured compensatory growth, but this came at a cost in terms of increased oxidative damage. Increased intake of antioxidants also resulted in changes in the activity concentration of enzymatic antioxidant defences and increased protein oxidative damage in both brain and blood. In addition, food deprivation caused increased protein oxidative damage in brain. Our findings show that the beneficial effects of dietary antioxidants on growth may be offset by hidden detrimental effects and that different early life events affect different components of oxidative status of a given tissue.

Keywords: Antioxidants, Aquaculture, Early life, Fish, Food restriction, Oxidative stress
1. Introduction

Positive selection of individuals capable of reaching an adult size quickly should be expected to sift out slowly growing individuals. Achieving an adult size as soon as possible should carry potential benefits to the individual, such as earlier time to sexual maturity or reduced predation risk (Dmitriew, 2011). Conversely to this prediction, animals do not grow at the maximum rate and much variation in growth strategies persists in both wild and laboratory animals, even when trophic resources are abundant (Blanckenhorn, 2000). The reason for this might lie with the costs of growing fast (Metcalfe and Monaghan, 2001). For example, rapid growth can increase metabolic demands (Criscuolo et al., 2008; Careau et al., 2013) and daily energy expenditure (Careau et al., 2013), or reduce body performance and lifespan (Lee et al., 2016). An additional mechanism through which growth may relate to self-maintenance is that increased cellular activity due to fast growth increases the production of reactive species, such as free radicals (Costantini, 2014; Smith et al., 2016). These unstable molecules, if not counteracted by antioxidants, oxidise proteins, lipids and nucleic acids, causing damage, and hence a state of oxidative stress (Halliwell and Gutteridge, 2015).

Recent correlative, experimental and meta-analytical work has suggested that oxidative stress might play a key role as a constraint on, and cost of, growth (Alonso-Alvarez et al., 2007; Nussey et al., 2009; Smith et al., 2016).

Oxidative costs of a rapid growth are particularly relevant when individuals accelerate growth to achieve the body size that is supposed to be normal for their life stage (known also as catch-up growth; Metcalfe and Monaghan, 2001). For example, abrupt reductions in food availability - common in wild populations - might lead young to slow down growth. When food availability becomes adequate again to meet the
nutritional requirements of a growing individual, juveniles will compensate by accelerating their growth. Although there are benefits in doing so, an acceleration of growth may also carry costs, such as a reduction in motor skills, sexual ornamentation, cognitive abilities or longevity (Metcalfe and Monaghan, 2001; Lee et al., 2016), but also in increased molecular oxidative damage (Smith et al., 2016).

Organisms may mitigate oxidative stress through an increased intake of antioxidants from diet. Animals normally ingest a cocktail of antioxidants and distribute them differently across tissues, depending on the chemical properties (lipophilic vs. hydrophilic) of the antioxidant and the specific requirements of a certain tissue (Surai, 2002; Halliwell and Gutteridge, 2015). Dietary antioxidants may be particularly important for growing individuals as the endogenous antioxidant machinery takes time to become fully mature (Surai, 2002; Costantini, 2014). In many vertebrate species, there is a greater reliance upon non-enzymatic antioxidants at earlier stages of development, such as those acquired from diet or from the mother (Surai, 2002; Costantini, 2014). It would therefore be expected that if an individual catches up while feeding on a diet rich in antioxidants, it might mitigate the oxidative costs of a fast growth. This simple prediction, however, does not consider that increases in the intake of a given antioxidant may also result in no net reduction in oxidative damage because any potential beneficial effects might be offset by compensatory reductions in other antioxidants. For example, long-term intake of a dietary antioxidant (vitamin C) in laboratory mice reduced expression of several genes linked to free radical scavenging, with no net effect on various oxidative damage metrics (Selman et al., 2006). A recent meta-analysis also showed that the effects of antioxidant supplementation on growth are
highly variable (from positive to negative), implying that effects of dietary antioxidants on oxidative status might be difficult to predict (Smith et al., 2016).

Another aspect that has received little attention is the impact that accelerated growth has on brain oxidative status and protein oxidative damage (Smith et al., 2016). This is surprising given that acceleration of growth may reduce cognitive abilities (Metcalfe and Monaghan, 2001), brain is a tissue particularly sensitive to oxidative stress (Barja, 2004) and protein turnover is altered during fast growth (Samuels and Baracos, 1995).

Compensatory growth is particularly common in fish at almost every stage of their life cycles because of their indeterminate growth patterns (Ali et al., 2003). Despite the expansion of the field of ecological oxidative stress, relatively little is known about the role of oxidative stress in mediating key life-history stages of fish (Birnie-Gauvin et al., 2017), such as those fish go through during growth and development.

In this study, we tested experimentally whether feeding on a diet rich in antioxidants enables young European seabass (Dicentrarchus labrax) to catch up after a period of food deprivation with negligible effects on the oxidative stress as compared to fish feeding on a normal diet (i.e., not enriched in antioxidants). To this end, we have measured metrics of oxidative damage and of antioxidant enzyme activity (i.e., proxy of enzyme concentration) in both blood and brain in juvenile fish that underwent a food deprivation period and then a refeeding protocol either with or without an enriched diet.

2. Materials and methods

2.1. Ethical note
All experiments complied with the Guidelines of the European Union Council and were approved by the Ethical Committee of the Tuscia University for the use of live animals (D.R. n. 677/16 and D.R. 644/17).

At the end of the food deprivation period, the loss of body mass in fish that were food-deprived was 11.4 and 12.6 % for the two experimental groups (mean ± se: prior food deprivation, 59.7 ± 5.1 g and 59.1 ± 5.1 g; end of food deprivation, 52.8 ± 5.2 g and 51.6 ± 5.2 g), respectively. Hence, it was below the threshold of 15 % recommended by the UK Home Office and the Canadian Council for Animal Care. The mortality rates calculated using sample sizes at the beginning of Phase 1 were similar among groups at the end of the experiment: 12.5 % (number of fish dead were 3), 25.0 % (number of fish dead were 7), 14.3 % (number of fish dead were 4) and 14.3 % (number of fish dead were 4) for no food deprivation/standard diet, no food deprivation/enriched diet, food deprivation/standard diet, and food deprivation/enriched diet, respectively.

2.2. Animals and husbandry

Juvenile European seabass (n = 112, one year of age) were obtained from a commercial hatchery (Cooperativa Orbetello Pesca Lagunare, Santa Liberata, Orbetello, Italy) located on the coast at about 40 km from the facility where the study took place. The fish were transported to the facility and randomly allocated into four fiberglass tanks (80 x 80 x 403 cm, all equipped with aeration system) in groups of similar size (26, 29, 29 and 28 individuals). Before being released into the tanks, each fish was massed and measured (mean ± SD, body mass: 64.5 ± 22.7 g; body length 18.9 ± 2.0 cm; body width 4.2 ± 0.5 cm, see below for method details). In each tank, fish were maintained in
a volume of 1,290 liters on an open circuit system with flow of 30 liters/minute
(replicating the conditions of the supplier hatchery) on natural light dark cycle and
illumination, as ample windows were present in the building. Being an open circuit
meant that each tank was subject to natural seasonal variations of chemical/physical
parameters: pH ranged between 8.0 and 8.2, temperature varied from 15°C in January to
23°C in August, while salinity ranged between 35 and 38 PSU; ammonia and nitrite
were below detectable levels, while nitrate was about 5 mg/L. The European seabass is
an euryhaline coastal species known to tolerate (at both juvenile and adult stage) a large
range of chemical-physical water parameters, including salinity and temperature (in the
order of 5-28°C) extremes (e.g. Dulger et al., 2012).

Tanks were cleaned daily to avoid accumulation of food remains. Fish were fed
by hand ad libitum once per day on the same commercial standard feed (company name
undisclosed) used by the supplier for 12 acclimation days. In order to make sure the
ration was ad libitum, we provided food until fish did not eat pellets anymore. The
residual pellets that deposited at the bottom of the tank were removed within around 60
minutes in order to minimise any potential confounding effects arising from very
bold/dominant individuals monopolising the access to the food. The feed composition
was as follows: crude proteins 48.0 %; crude fats and oils 22.0 %; crude fibre 3.0 %;
crude ash 6.0 %; vitamin E 150 mg/Kg.

2.3. Experimental procedure and sampling
Phase 1, food deprivation: after the acclimation period, the fish of two tanks started a
food deprivation protocol that lasted 81 days, while the control fish in the other two
tanks continued to be fed daily on the same commercial feed as previously (Fig. 1). All
fish were massed and measured on the day Phase 1 was started; the sample size was 108 because four fish died before the start of Phase 1.

Phase 2, re-feeding and enriched diet: as the food deprivation protocol stopped, the pool of animals was re-fed as follows: two groups were fed daily on the standard feed, and the other two groups were fed by hand *ad libitum* once per day on a feed enriched with vitamin E (feed composition: crude proteins 48.0 %; crude fats and oils 20.0 %; crude fibre 2.0 %; crude ash 6.0 %; vitamin E 600 mg/Kg). In this way any effect of the enriched diet on growth and oxidative status could be tested on both the food restricted and the control fish (Fig. 1). This phase lasted until the final sampling, which occurred 180 days later.

In order to quantify the growth, fish were caught with hand nets by two operators, massed (Sartorius TE 612) and measured with a caliper for length and body width (excluding dorsal and ventral fins) by a third operator once at the end of phase 1 and three times during phase 2 (Fig. 1). At the final sampling, the animals were bled after a mild anesthesia and were straightaway euthanized with an overdose of 2-phenoxiethanol (8 ml/l). Soon after euthanisation, the brain was collected from each individual. Tubes containing blood samples were spun to separate plasma from red blood cells. All samples were stored at -80°C and later transported in dry ice to the laboratories of the University of Antwerp for the biochemical analyses.

2.4. Laboratory analyses

We randomly selected samples of blood and brain of 10 individuals per experimental group. Whole brain was homogenised in cold PBS (supplemented with 20% (v/v) of glycerol and with 0.2 mM of phenylmethylosulfonyl fluoride as an inhibitor of proteases)
using a pestle. Samples were then sonicated for 10 minutes and then centrifuged for 10 minutes at 15,000 rpm. The supernatant was taken and split into different tubes, which were stored at -80°C for later analyses. Haemolysates were centrifuged to separate cell membranes from the supernatant, which were used for the analyses straightaway. The d-ROMs assay (reactive oxygen metabolites; Diacron International, Grosseto, Italy) was used to measure plasma oxidative damage metabolites (e.g., organic hydroperoxides) that are generated early in the oxidative cascade. Values were expressed as mM H$_2$O$_2$ equivalents. Protein carbonyls (marker of oxidative damage to proteins) were measured in plasma and brain homogenates using the Protein Carbonyl Colorimetric assay (Cayman Chemical Company, Ann Arbor, MI, USA). The assay is based on the protocol of Levine et al. (1990). The concentration of protein carbonyls has been expressed as nmoles/mg proteins. The Ransod assay (Randox Laboratories, Crumlin, UK) was used to quantify the activity of the enzyme superoxide dismutase (SOD) in both red blood cells and brain. The activity of SOD has been expressed as Units/mg proteins. The Ransel assay (Randox Laboratories, Crumlin, UK) was used to quantify the activity of the enzyme glutathione peroxidase (GPX) in both red blood cells and brain. The activity of GPX has been expressed as Units/mg proteins. The assay is based on the original method of Paglia and Valentine (1967). Protein concentration in the haemolysate or homogenate was measured using the Bradford protein assay (Bio-Rad Laboratories, Hercules, CA, USA).

2.5. Statistical analyses

Statistical analyses were performed using SPSS Version 23. General Linear Models were used to test the effect of our experimental manipulation on growth. Three main
factors and their interactions were included in the model. The main factors were pre-
refeeding group (fed or food deprived), refeeding group (food enriched or not enriched
with antioxidants) and sampling day (three levels). As a response variable (body size
index), we used the first principal component (PC1, 95.8 % of the total variance)
extracted from a principal component analysis (PCA) on body mass, body length and
body width. The loadings (expressed as Pearson correlation coefficients) of these three
body measurements on the PC1 were as follows: 0.984 for body mass, 0.984 for body
length and 0.969 for body width. Analyses were repeated including body length or body
size (PC1 of a PCA on body mass and body width) as response variables, respectively.
General Linear Models were also used to test the effect of our experimental
manipulation on each metric of oxidative status. Each model included two main factors
(pre-refeeding group and refeeding group) and their interaction. To meet normality of
residuals and homogeneity of variances, plasma reactive oxygen metabolites and brain
superoxide dismutase were both square-root transformed, while brain protein carbonyls
were log-transformed. Transformations were chosen according to the fitting of the
models using values of Akaike Information Criterion as a reference. Post-hoc tests were
used to explore further any significant interaction across logical sets of means. When
non-significant, interactions were removed from the models. Outcomes of full and
reduced models coincided when there was not backward elimination because the
interaction/s was/were significant.

3. Results

The difference in body size between control and treated fish depended on whether fish
were re-fed on a normal or enriched diet (three-way interaction: p = 0.047). To
investigate this further, the effect of the experimental manipulation on body size was
tested separately for fish re-fed on a normal or enriched diet, using similar linear models
as explained above. Fish re-fed on a normal diet did not catch-up, resulting in a smaller
body size than those fish that were not food-deprived (Fig. 2, Table 1). Conversely, fish
re-fed on an enriched diet were capable to catch-up, resulting in a body size as that of
fish that were not food-deprived (Fig. 2, Table 1). Similar results were obtained if body
size was extracted from a PCA calculated on body mass and body width (three-way
interaction: $p = 0.012$). As for body length, fish that were food-deprived were shorter
than fish that were not food-deprived at the end of the food deprivation period, while
they were of similar body length at the end of the experiment (Fig. 3; Table 1). The
effect of food deprivation was not influenced by the quality of diet given after the food
depprivation period (Table 1).

The difference in plasma reactive oxygen metabolites between control and
treated fish depended on whether fish were re-fed on a normal or enriched diet. Fish that
were fed on an enriched diet after food deprivation had significantly higher plasma
reactive oxygen metabolites than all other experimental groups (Fig. 4, Table 1).

Plasma protein carbonyls were higher in fish that were either fed continuously or
re-fed on an enriched diet than those either fed continuously or re-fed on a normal diet
(Fig. 5, Table 1). Irrespective of the quality of re-feeding diet, brain protein carbonyls
were higher in fish that were food-deprived than in fish that were not (Fig. 6, Table 1).

The activity of glutathione peroxidase in red blood cells was higher in fish that
were either fed continuously or re-fed on an enriched diet than those either fed
continuously or re-fed on a normal diet (Fig. 5, Table 1). The activity of brain
glutathione peroxidase was unaffected by the experimental manipulation in both phases (Table 1).

The activity of superoxide dismutase in both red blood cells and brain was lower in fish that were either fed continuously or re-fed on an enriched diet than those either fed continuously or re-fed on a normal diet (Fig. 5, Table 1).

4. Discussion

The results of this study show that a diet rich in vitamin E may favour compensatory growth after a prolonged period of fasting, but this comes at a cost in terms of increased oxidative damage. As expected, our food deprivation regime slowed down growth of fish. A re-feeding period of six months enabled fish to compensate through a faster growth, but the compensatory growth was mainly evident for mass/condition, while there was not compensation for body length (proxy of skeletal size). There are at least two possible explanations. Previous studies on fish have shown that (i) growth compensation may be limited in some species to restoring lost tissues/energy reserves, with no acceleration of skeletal growth (Alvarez and Nicieza, 2005), or (ii) growth compensation of body length may take longer than body mass/condition to be evident (e.g., Johnsson and Bohlin, 2006).

The compensatory growth occurred only in those fish that were re-fed with a diet enriched with vitamin E, highlighting the importance of this nutrient for fish growth. Vitamin E may interact with many metabolic and cellular pathways important for growth, such as lipid transport by lipoproteins, vitamin C synthesis, function of erythrocytes, immunity or metabolism of other nutrients (Hamre, 2011). Previous work on juvenile gilthead seabreams (Sparus aurata) also showed that increased intake of
vitamin E can reduce skeletal anomalies, favouring growth (Izquierdo et al., 2013). Vitamin E promotes membrane repair and such repair ability is particularly important when cells are being exposed to an oxidant challenge (Howard et al., 2011; Labazi et al., 2015). Moreover, vitamin E may reduce haemolysis of red blood cells through a chain-breaking activity, i.e., able to scavenge free radical by hydrogen donation (Niki et al., 1988; Niki, 2014). Although vitamin E has antioxidant properties, those fish that made a compensatory growth had higher levels of plasma reactive oxygen metabolites than those fish that did not make a compensatory growth. High metabolic activity typical of compensatory growth might have caused a high production of oxygen metabolites. Conversely, protein oxidative damage was not increased in fish that made a compensatory growth. A reason for this might lie with a higher protection of proteins against oxidation. Reactive oxygen metabolites as measured in our work come from early oxidation of biomolecules, and their accumulation is prevented by the activity of the glutathione peroxidase (Halliwell and Gutteridge, 2015). On the other hand, protein carbonylation is mostly irreversible damage, thus avoidance of carbonylation might have been prioritised and traded-off against other kinds of damage.

The higher production of reactive oxygen metabolites in those fish that made a compensatory growth might have also been favoured by the impact that vitamin E intake had on antioxidant enzymes. Previous work on fish showed that vitamin E may reduce the expression of superoxide dismutase gene and increase the expression of the glutathione peroxidase gene, respectively (Izquierdo et al., 2013). These results on expression of enzymatic antioxidant genes are in agreement with our results on the activity of superoxide dismutase and glutathione peroxidase. Superoxide dismutase catalyses the dismutation of the superoxide anion generated by mitochondria into
oxygen and hydrogen peroxide; glutathione peroxidase reduces hydrogen peroxide and organic hydroperoxides into water and alcohols, respectively (Halliwell and Gutteridge, 2015). Hence, lower activity of superoxide dismutase might have resulted in reduced protection against the peroxidative action of superoxide anion, which may increase generation of organic hydroperoxides (Halliwell and Gutteridge, 2015). Upregulation of glutathione peroxidase might have been needed to detoxify cells from organic hydroperoxides, which may propagate the oxidative cascade (Halliwell and Gutteridge, 2015). Fish feeding on the diet enriched with vitamin E had actually higher protein oxidative damage than those feeding on a normal diet.

Although vitamin E has antioxidant properties, any beneficial effects of having large amounts of vitamin E available do not necessarily come through its antioxidant properties. Selman et al. (2008) showed that supplementation of vitamin E significantly increased median lifespan of mice by 15% relative to non-supplemented controls and also increased maximum lifespan. Lymphocyte and hepatocyte oxidative DNA damage and hepatic lipid peroxidation were, however, unaffected by supplementation. Using a cDNA macroarray, Selman et al. (2008) showed that genes associated with xenobiotic metabolism were significantly upregulated in the livers of mice, suggesting that vitamin E is initially metabolized by cytochrome P450s in a manner similar to xenobiotics. The absence of any significant effect on oxidative damage suggests that the lifespan extension observed in mice was not due to the antioxidant properties of vitamin E.

In various invertebrates (e.g., molluscs, crustaceans) and fish, data on development of antioxidant defences suggested that dietary and other low molecular weight antioxidants are predominant in the earliest stages of development, while antioxidant enzymes become predominant later in life (Rudneva, 1999; Surai, 2002;
Fontagné et al., 2008). We do not know if any beneficial effects of dietary antioxidants could have been stronger at earlier stages of development. This is an open question that deserves further work to understand at which developmental stage intake of dietary antioxidants might be more beneficial. It will also be important to assess the amount of tocopherols that can be retained in tissues and which forms of tocopherols and of their metabolites have a stronger impact on the antioxidant machinery during development (Brigelius-Flohé and Traber, 1999; Surai, 2002).

Our results also show that experiencing a period of food deprivation may have long-term effects on the protein oxidative damage in brain. Irrespective of the quality of re-feeding diet, all the fish that underwent food restriction had higher oxidative damage in brain later in life than those that did not. Restriction of intake of specific nutrients in early life may have pervasive, long-lasting effects on oxidative status. For example, rats subjected to prenatal and postnatal protein malnutrition had increased oxidative damage to lipids and proteins in brain (Feoli et al., 2006). Zebra finches (Taeniopygia guttata) that experienced a short period of low quality nutrition during the nestling period had a twofold reduction in plasma levels of carotenoids, vitamins A and E in adulthood (Blount et al., 2003). In fish, starvation may elicit profound changes in brain energy metabolism in a way similar to that demonstrated in other vertebrates like mammals. For example, when the fasting period is prolonged and glucose becomes less available with time, the brain obtains a portion of its energy from ketone bodies (Hasselbalch et al., 1994; Campbell and Farrel, 2006), which are important generators of protein carbonylation (Halliwell and Gutteridge, 2015). Moreover, during prolonged fasting, animals face increased protein catabolism and body mass loss (McCue, 2010, 2012), which might favour accumulation of protein carbonyls.
In conclusion, our work provides evidence that any beneficial effects of dietary antioxidants on growth might be offset by an impaired capability of regulating the oxidative status. If investing more into compensatory growth is more advantageous than into self-maintenance and mechanisms favouring lifespan likely depends on the environmental conditions young develop in, such as predation risk or food availability. Work on oxidative stress resistance under this life-history framework would help to clarify its role as a mediator of some early life tradeoffs across different environments. Our results also point for a role of oxidative stress as a mechanism linking the long-term negative effects of early life poor nutrition on brain function.

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Competing interests

No competing interests.

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Table 1. Outcomes of full and reduced statistical models. FD = Food Deprivation; POSTFD = diet after the food deprivation. The reduced models were obtained after backward elimination of non-significant interactions from the full models, starting from the three-way interaction. Outcomes of full and reduced models coincided when there was not backward elimination because the interaction/s was/were significant. Markers of oxidative status were analysed in a subsample of 40 individuals (10 for each group).

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<tr>
<td>Treatment group FD x</td>
<td>Treatment group POSTFD</td>
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Figure captions

Figure 1. Experimental setting and timeline.

Figure 2. Food deprivation caused a slowdown of growth as suggested by the smaller body size compared to fish that were not food-deprived. Refeeding caused accelerated growth only in those fish that were fed on a diet richer in vitamin E.

Figure 3. Food deprivation caused a slowdown of skeletal growth as suggested by the smaller body length of food-deprived fish compared to fish that were not food-deprived soon after the food deprivation period (Tukey test, p<0.01) and at the end of the re-feeding experiment (Tukey test, p<0.01).

Figure 4. Fish that made a compensatory growth (black square) had higher plasma oxidative damage (expressed as reactive oxygen metabolites).

Figure 5. Fish fed on the diet richer in vitamin E had higher plasma protein oxidative damage (protein carbonyls) and activity of glutathione peroxidase in red blood cells (RBS) and lower activity of superoxide dismutase in both red blood cells and brain.

Figure 6. Fish that experienced food deprivation had higher brain protein oxidative damage (protein carbonyls) than those fish that were fed ad libitum.
Figure 1

- Sampling of body mass, body length and body width
- Sampling of body mass, body length, body width and tissues
Body Size (PC1)

- NO FOOD DEPRIVATION - STANDARD DIET
- FOOD DEPRIVATION - STANDARD DIET
- NO FOOD DEPRIVATION - ENRICHED DIET
- FOOD DEPRIVATION - ENRICHED DIET

BEFORE FOOD DEPRIVATION
AFTER FOOD DEPRIVATION
END OF EXPERIMENT

Figure 2
Figure 3

Body length (cm)

NO FOOD DEPRIVATION
FOOD DEPRIVATION

BEFORE FOOD DEPRIVATION  AFTER FOOD DEPRIVATION  END OF EXPERIMENT
Figure 4

SQRT Reactive Oxygen Metabolites (mol l$^{-1}$ H$_2$O$_2$ equivalents)

- NO FOOD DEPRIVATION - STANDARD DIET
- FOOD DEPRIVATION - STANDARD DIET
- NO FOOD DEPRIVATION - ENRICHED DIET
- FOOD DEPRIVATION - ENRICHED DIET
Figure 5

- Plasma Protein Carbonyls (nmol mg⁻¹ proteins)
- RBC Glutathione Peroxidase (U/mg-proteins)
- RBC Superoxide Dismutase (U/mg-proteins)
- Brain Superoxide Dismutase (U/mg-proteins)

STANDARD DIET
ENRICHED DIET
Figure 6

Log (Brain Protein Carbonyls + 1)
(nmol mg⁻¹ proteins)

○ NO FOOD DEPRIVATION
■ FOOD DEPRIVATION