Mommy Fishues

 $\label{eq:continuous} \mbox{A Thesis}$ $\mbox{Presented to}$ $\mbox{The Established Interdisciplinary Committee for Neuroscience}$ $\mbox{Reed College}$

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Marissa Breanne Borrego

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Suzy C. P. Renn

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Any situation in which some individuals prevent others from engaging in the process of inquiry is one of violence.

> Paulo Freire Pedagogy of the Oppressed

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Preface

Science has a history as an oppressive institution. That being said, I think that science also has the ability to liberate individuals. I hope that at the least this thesis makes one think of how plastic we are to our day-to-day experiences.

List of Abbreviations

 11β -HSD2 11β -hydroxysteroid dehydrogenase type 2

ACTH Adrenocorticotropin hormone
CRH Corticotropin-releasing hormone
CPP Conditioned place preference

GR Glucocorticoid receptor

GRE Glucocorticoid response element

HLG High licking & grooming

HPA Hypothalamic-pituitary-adrenal

LLG Low licking & groomingPCR Polymerase chain reactionPVN Pareventricular nucleus

qPCR Quantitative polymerase chain reaction

Table of Contents

Introd	uction	1
Chapte	er 1: S	tress
1.1	The H	ypothalamic-Pituitary-Adrenal Axis
	1.1.1	Activation of the HPA Axis
	1.1.2	Glucocorticoid Receptors
	1.1.3	Regulation of the HPA Axis
1.2	Stress	and Development
	1.2.1	Protection Against Early Life Stress
	1.2.2	The Effects of Prenatal Stress on Development
	1.2.3	Maternal Care and Stress in Rats
	1.2.4	Early Social Environment and Stress in Cichlids
	1.2.5	Astatotilapia burtoni
	1.2.6	Current Investigation
Chapte	er 2: T	The Experiment
2.1		$\mathrm{d}\mathrm{s}$
	2.1.1	The Fish
	2.1.2	Behavioral Tests
	2.1.3	Gene Expression Assay
2.2	Result	s
	2.2.1	Size
	2.2.2	Boldness
	2.2.3	Aggression
	2.2.4	Glucocorticoid Receptor Expression
2.3	What	Does this Mean?
2.4		graphies
Refere	nces	21

List of Tables

1.1	Summary of Meaney's work comparing high licking and grooming and	
	low licking and grooming pup phenotypes	10
1.2	Summary of Taborsky's work comparing <i>N. pulcher</i> fry raised in the	
	presence and absence of adults and adolescents	11
2.1	Genes and corresponding primer sequences used for qPCR	19

List of Figures

1.1	Signaling cascade of the HPA axis	4
1.2	Glucocorticoid receptor mechanism of action	6
2.1	Experimental conditions	15
2.2	Behavioral paradigms	17
2.3	Timeline of experiment	18
2.4	Primer design	18
2.5	The central dogma of molecular biology including reverse transcription	19

Abstract

The preface pretty much says it all.

Dedication

And in our hearts How beautiful the flames that will flare up in a ring

Chika Sigawa "Mountain Range"

For Langston.

Introduction

A defining feature of living organisms is that they are able to respond to stimuli in their environment. In other words, they behave. Each behavior requires a stimulus, or multiple stimuli, that triggers a chain reaction of internal responses, changing how an organism exists in its environment. In understanding why an animal responds to a stimulus in the way that it does, there are two places to start. One can look to the organism's genotype *i.e.*, was this behavior inherited genetically from its parents? Or one can look to the organism's upbringing *i.e.*, was this behavior learned in response to the environment. Traditionally, these two possibilities have been thought of as separate and exclusive, as in the phrase "nature vs nurture".

The dichotomy of nature and nurture as we know it today has its unfortunate beginnings in the field of eugenics. The phrase was popularized by the father of eugenics, Francis Galton, in the late 19th century in an effort to understand if human "ability" was heritable. He defined nature as "all that a man brings with himself into the world" and nurture as "every influence from without that affects him after his birth" (Galton, 1874). While there was not yet a concept of DNA, both Darwin's theory of evolution and Mendel's inheritence experiments were in circulation. The interest in nature vs nurture remained within developmental psychology until late in the 20th century when behavioral and developmental neurosciences were popularized.

In the early and mid 20th century, the fields of animal behavior and genetics were being revolutionized in ways that would ultimately contribute to the modern debate of nature and nurture (Krubitzer and Kahn, 2003). In the 1930's a pioneering behavioral scientist by the name of Nikolaas Tinbergen began studying behaviors holistically, as a product of individual experience and evolution. He was interested in creating a scientifically rigorous way by which to observe and comment on behavior. What emerged was the modern field of ethology and a set of four categories to study a behavior through: causation (mechanism), survival value (adaptation), ontogeny, and evolution (Tinbergen, 2005). Tinbergen's four questions were important in examining a single behavior as a product of an individual's experiences and that individual's

2 Introduction

lineage. That being said, there was still not that much known about molecular biology and its role in behavior.

Abstract concepts of DNA and RNA as a heritable molecule had been proposed by the early 20th century in response to heritability studies (Koltzoff, 1934; Hershey and Chase, 1952), but it wasn't until Francis Crick and James Watson published a study in 1953 on the structure of DNA (notably, the study relied heavily on prior work by Rosalind Franklin) that the field of modern genetics really began (Watson and Crick, 1953). Using information about base pairs and amino acids published by other labs at the time, Crick proposed the central dogma of genetics in 1955. This crucial concept states that DNA is translated into RNA, which is then transcribed into amino acids that are linked together to form proteins (see Figure 2.4).

The last big step in getting to our current concept of nature and nurture was the popularization of epigenetics. Epigenetics, in short (this concept will be revisited in Chapter 1), refers to the factors that change the ability of DNA to be transcribed, contributing to changes in gene expression. Much of modern behavioral sciences is aimed at understanding how the environment influences an organisms epigenome.

Because we now understand gene expression is often altered by the environment, our notion of nature vs nurture becomes rather arbitrary. Behaviors can instead be thought of as an intertwining of nature and nurture. Rather than understanding the ratio of environmental to genetic influence on a behavior, we can instead examine how certain genotypes make an organism more vulnerable to environmental influences or how the environment influences the ways in which the genome is utilized. This research is crucial because it implies that, as animals, our behavior can be drastically altered by our environment. We must strive to understand how the spaces we create impact one another and to interrogate the ways in which we judge people's ability to control their actions.

Put simply, this thesis will examine how maternal care (a stimulus) affects stress response (a behavior) in a mouth-brooding fish. I will begin by explaining the physiological components of the stress response and how this can be changed by the environment. Then I will give a brief summary of what is already known about maternal care's influence on behavior, before diving into the novel research I conducted. I will conclude with the implications of my study and future directions for this line of research.

Chapter 1

Stress

1.1 The Hypothalamic-Pituitary-Adrenal Axis

If you have made it this far in life, you have at some point felt *stressed*. Stress can be defined as the body's *response to* and *recovery from* a threat that disrupts homeostatis (van Bodegom et al., 2017). An important aspect of the stress response is the production and mobilization of energy. This is made possible through the hypothalamic-pituitary-adrenal (HPA) axis, which functions to produce glucocorticoids. As the name suggests, glucocorticoids play a role in the metabolism of glucose, the body's main source of energy. The following section will introduce the the mechanisms involved in HPA axis activation.

1.1.1 Activation of the HPA Axis

The activation of the HPA axis begins with stimulation of the hypothalamus by other brain areas. In the presense of an immediate stressor, brain regions associated with maintaining homeostasis trigger the axis. Take for example the response to a painful stimulus. Pain is sensed by nociceptors in the peripheral nervous system and cause afferent signaling to norepinephrinergic neurons in the hind brain. These hindbrain neurons can in turn stimulate the hypothalamic neurons involved in the HPA axis. It is also possible to activate the HPA axis as an anticipatory response. If an animal has been conditioned to associate a given smell with a predator, then in the presence of that smell alone the animal may trigger the HPA axis in anticipation of the danger. This requires polysynaptic signaling from limbic structures involved in learning and fear such as the hippocampus and amygdala. The hippocampus excites the axis through glutamatergic interneurons, whereas it is hypothesized that much of

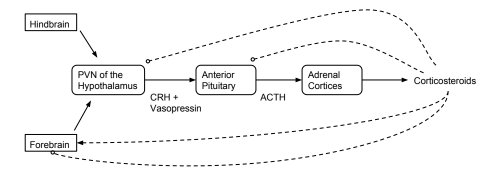


Figure 1.1: Signaling cascade of the HPA axis.

As a response to stress, forebrain or hindbrain projections to the hypothalamus can begin the HPA axis signalling cascade. The paraventricular nucleus (PVN) of the hypothalamus secretes corticotropin-releasing hormone (CRH) and vasopressin into the anterior pituitary. This causes the anterior pituitary to release adrenocorticotropin hormone (ACTH) into the bloodstream. ACTH reaches the arenal cortices of the adrenal glands, leading to the production of steroid hormones including corticosteroids. While corticosteroids have many targets and regulatory effects, they can also have positive feedback effects (dashed arrowhead) and negative feedback effects (dashed open circle) for the HPA axis itself.

the excitatory amygdalar signaling works through disinhibition In both the immediate and anticipated cases, the activation of neurons within the hypothalamus leads to a sterotyped cascade of signalling.

The hypothalamus is a region of the midbrain known for its role in maintaining allostasis through its involvement in stress, appetite, circadian rhythms, and sexual behavior. In response to a stressor, the paraventricular nucleus (PVN) of the hypothalamus secretes corticotropin-releasing hormone (CRH) and vasopressin, which bind to receptors in the pituitary gland. The pituitary gland is directly ventral to the hypothalamus and is a main regulator of hormone release. The binding of CRH to CRF₁ receptors in the anterior pituitary leads to the secretion of adrenocorticotropin hormone (ACTH). This excitatory interaction can be potentiated by vasopressin, though vasopressin alone is not enough to produce an effect. ACTH enters the blood stream and travels to the adrenal cortices, which are the dorsal regions of the adrenal glands. ACTH binds to melanocortin 2 receptors, which increases the synthesis of cholesterol. Cholesterol is then transported to the outer mitochondrial matrix where the steroidogenic pathway begins. A major end product of this pathway and thus of the HPA axis is corticosteroids.

1.1.2 Glucocorticoid Receptors

Corticosteroids are hormone peptides that can bind to glucocorticoid receptors (GRs) and mineralocorticoid receptors. The term glucocorticoid refers to corticosteroids that are able to bind to GRs. After being released by the adrenal glands, glucocorticoids travel through the blood stream, pass through the blood-brain-barrier, and freely diffuse into the cytoplasm of neurons. Unbound GRs reside in the cytoplasm as part of a larger protein complex. When a GR is bound to it goes through a confirmational change and sheds the protein complex. It is then transported to the nucleus where it dimerizes with another GR. The homodimer can then interact with other proteins, ultimately leading to the binding of the complex to a glucocorticoid response element (GRE) on the genome. These GREs are often found in the promotor region of their target genes and recruit transcription factors to suppress or enhance transcription of the target gene (Busada and Cidlowski; Herman et al., 2005). Interestingly, GRs can also localize to the mitochondria, where they can alter transcriptional elements of mitochondrial DNA. Due to mitochondria's role in steroid and energy production (mitoch), this finding suggests an important long-term regulatory role of glucocorticoids. Ligand binding to GRs can also have nongenomic consequences such as kinase activation, though these pathways are not well understood (Samarasinghe et al., 2011).

In mammals, there are eight known transcriptional isoforms of the GR gene (Saif et al., 2015) and thirteen different post-translational modification sites of the GR protein (Oakley and Cidlowski, 2013). These differences in protein structure alter the cellular function of the GR subtypes (Lu et al., 2007). A genome duplication event happened in the evolution of teleosts, causing them to have two GR paralogues: GR1 and GR2. The genetic sequences are highly similar to each other as well as to the GR genes of other species (Greenwood et al., 2003). Both GR1 and GR2 are expressed in corticosteroid responsive regions, suggesting that they both maintain signaling functionality.

1.1.3 Regulation of the HPA Axis

Activation of the HPA axis leads to situationally different levels and duration of glucocorticoid release. Inflammatory stressors often lead to prolonged stress responses, as the injury requires sustained energy to repair. Psychological stressors, in contrast, tend to lead to acute responses because there is no tissue repair or immune response required (Herman et al., 2016). Because the stress response is energy intensive, it is important that an organism responds appropriately to threatening situations.

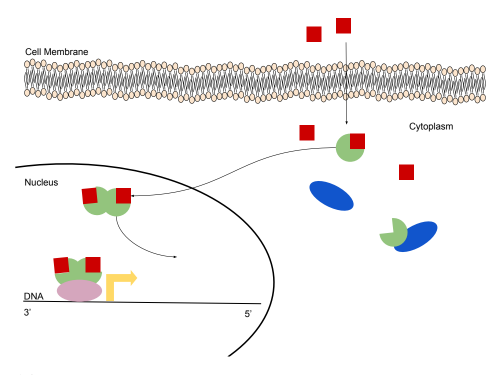


Figure 1.2: Glucocorticoid receptor mechanism of action.

Unbound glucocorticoid receptors (green) reside in the cytoplasm of cells as a part of a larger protein complex (blue). Corticosteroids (red) can freely diffuse through the cell membrane and bind to glucocorticoid receptors. Glucorticoid receptors then undergoe a confirmational change, shedding the protein complex, and move into the nucleus. There they form homodimers and bind to glucocorticoid response elements (pink) on the genome. These elements are downstream of promotor regions (yellow) and can alter the transcription of nearby genes.

The HPA axis has built in negative feedback loops to ensure tight regulation. GR's play an inhibitory role in the activation of the HPA axis. They are abundantly expressed within the PVN. Upon activation, they cause endocannabinoid synthesis and release that are able to inhibit glucocorticoid receptors that target CRH neurons. Long term exposure to corticosteroids has also been shown to reduce pituitary ACTH release (Keller-Wood, 2015). In addition to regulation within the axis, GRs in the hippocampus and prefrontal cortex are able to inhibit HPA axis activity via GABAergic interneurons.

The HPA axis has the ability to adjust to chronic stress. There is an important distinction to be made by the body between long-term stressors that continuously pose a threat to an organism and long-term stressors that don't actually pose a threat to an organism. Take for example two deer that live in environments coinhabited by humans. The first lives in an area that is frequented by hunters. This deer has to induce a full stress response every time it encounters a human, else the animal will quickly be killed. Now take for example a deer that lives in a zoo. It would surely be a waste of this deer's energy if it were to enter a stressed state every time it encountered a human. While it is adaptive for the former deer to develop a heightened responsiveness to humans, it is adaptive for the latter deer to become desensitized to humans.

A long-term change to the physiology of the HPA axis can occur via epigenetic modification. In other words, the genes encoding proteins necessary for the HPA axis can be made more or less likely to be transcribed as a result of chemical changes to structural elements of the DNA. Epigentic changes are heritable from parent to daughter cell even though they do not alter the actual DNA sequence, making them an important aspect of early development when cells are rapidly proliferating.

1.2 Stress and Development

The fact that chronic stress is often unhealthy is quite intuitive. When an organism is forced to expend energy on immediate survival, it must forego less pressing, but very important processes like eating, sleeping, reproducing, and learning. The following section will examine some of the biological underpinnings of the maladaptive nature of chronic stress.

1.2.1 Protection Against Early Life Stress

There exists a highly conserved protection against early life stress in both mammals and teleosts. In both cases, mothers secrete 11β -hydroxysteroid dehydrogenase type 2 (11β -HSD2) into the prenatal environment. This hormone rapidly deactivates corticosteroids, effectively inhibiting the stress response of embryos (van Bodegom et al., 2017; Faught et al., 2016). As newborns, there is a stress hyporesponsiveness period characterized by a decrease in circulating ACTH and corticosteroids, as well as an overall decrease in responsiveness to stressors (van Bodegom et al., 2017; Barry et al., 1995).

Importantly, both of these defenses can be altered by a highly stressful environment. Repeated maternal exposure to stress decreases 11β -HSD2, increasing prenatal corticosteroid exposure. Additionally maternal separation is associated with a shortened stress hyporesponsiveness period in mammals (no similar study has been done in fish). These findings suggest that the stress response is plastic to early life experience. This is evolutionarily favorible in that it allows the animal to adapt to its environment. The match/mismatch hypothesis suggests that organisms experiencing adversity early in life will most likely face similar levels of adversity later in life and should adjust their stress response to reflect that. With this hypothesis comes the idea that a mismatch in early environment and adult environment would have adverse consequences for the organism (Gluckman et al., 2007).

1.2.2 The Effects of Prenatal Stress on Development

Glucocorticoid receptors play an important role in fetal development. While insufficient levels of glucocorticoids are sometimes fatal, causing undeveloped organs, excess circulating glucorticoids are also potentially maladaptive, leading to developmental reprogramming (Busada and Cidlowski). The offspring of pregnant mice treated with synthetic glucocorticoids have delayed maturation of neurons and glia as well as delayed vascularization of the brain (Gravanis and Mellon, 2011). Further, prenatal stress exposure is correlated with decreased dendritic spine density in the cingulate gyrus and orbitofrontal cortex (Murmu et al., 2006). These data suggest that prenatal stress alters the physiology and connectome of the developing brain.

The ability of offspring to learn and form memories is altered by prenatal stress exposure. Compared to controls, offspring of stressed mothers have decreased fear learning in a passive avoidance behavioral paradigm (Sofiabadi et al., 2018). Older rats have impaired spatial memory in the Y-maze and working memory in the radial

arm maze when they were exposed to prenatal stress (Vallée et al., 1999). Rats also display a corresponding decrease in CAMKII and CREB mRNA expression in the hippocampus under these conditions (Sun et al., 2017).

Animals exposed to prenatal stress are also more susceptible to addictive behavior. Rats with mothers exposed to stress have increased nicotine conditioned placed preference (CPP) as well as increased dopamine D₂ receptor gene expression in nucleus accumbens (Said et al., 2015). Prenatal stress has also been shown to increase CPP in response to benzodiazepines (Lakehayli et al., 2015), cocaine (Pastor et al., 2018), and morphine (Vey et al., 2016) just to name a few.

Prenatal stress is a predictor of psychiatric disease in adults. Rodents exposed to prenatal stress show increased anxiety, depressive, and schizophrenic-like behavior compared to offspring of non-stressed dams (Weinstock, 2017).

1.2.3 Maternal Care and Stress in Rats

Stress in the postnatal environment can also be influential to an organism's development. The lab of Michael Meaney has done years of groundbreaking work on how maternal care alters the stress response in rats. Rat mothers exhibit consistent differences in the time spent licking and grooming their young during their first week of life (Meaney et al., 1996). This difference takes place during a critical period of the rats neural development. As a result, pups reared by high licking and grooming (HLG) mothers and low licking and grooming (LLG) mothers have distinct phenotypes and epigenomes (Weaver et al., 2004).

In 1997, Meaney's lab examined how circulating stress hormones differed in pups reared by HLG and LLG mothers (Liu et al., 2000). HLG pups had reduced circulating levels of ACTH and corticosterone in response to restraint stress. Additionally, HLG pups appeared to have enhanced regulatory feedback in stressful situations, as they surpressed ACTH to a greater extent after being pre-treated with corticosterone (the murine equivalent of cortisol). HLG pups also developed higher GR expression in the hippocampus as adults. In a complementary study, the lab found a distinct behavioral phenotype between the two groups (Caldji et al., 1998). Rats reared by HLG dams exhibited more exploratory behavior, as measured by an open field paradigm, compared to those reared by LLG dams. Additionally, LLG pups exhibited a longer latency to start eating when placed in a novel environment compared to HLG pups. Meaney cross-fostered pups from HLG and LLG mothers. As a result, pups born to LLG dams, but reared by HLG dams had a similar phenotype to those born

to and reared by HLG dams (Francis et al., 1999). These findings indicated that maternal care can influence offsprings' responses to stress as adults.

In 2004, Meaney published a paper on the epigenetics of the above discoveries (Weaver et al., 2004). He found that the epigenetic state of the GR promoter gene was altered by maternal licking and grooming. This difference in methylation state was contingent on the rearing, not the biological, mother. GR receptor gene methylation was decreased and acetylation increased in HLG rats, consistent with earlier studies. Taken together, Meaney's work demonstates that rats reared by LLG dams have a hypersensitive stress response, characterized by increased circulating stress hormones, decreased hippocampal GR receptors, and decreased exploratory behavior.

Table 1.1: Summary of Meaney's work comparing high licking and grooming and low licking and grooming pup phenotypes.

Phenotype	Pups Reared by HGL Dams	Pups Reared by LLG Dams
Circulating stress hormones	Low	High
Habituation to Corticosterone	High	Low
Hippocampal GR expression	High	Low
Anxiety-like behavior	Low	High
GR gene methylation	Low	High
GR gene acetylation	High	Low

1.2.4 Early Social Environment and Stress in Cichlids

Barbara Taborsky has examined how early social environment influences cichlid stress response. Most of Taborsky's work is with *Neolamprologus pulcher*, a highly social cichlid species that lives in family units and collectively raises offspring. Immature fish help to keep eggs clean and well-oxygenated while adults defend the eggs against predators and conspecifics (Arnold and Taborsky, 2010).

Much of Taborsky's work has focused on how early life social experience affects social behavior and stress response in adults. Fish were divided into three groups: those raised with adults and immature helpers, those raised with just helpers, and those raised with neither helpers nor adults. In the following studies, fry raised in the presence of just helpers and fry raised in the presence of helpers and adults had the same trends. Taborsky found that *N. pulcher* fry raised in the absense of adults and helpers had decreased social competency, showing energetically costly levels of aggression in territory disputes (Arnold and Taborsky, 2010). Fish raised with adults

had decreased whole-brain GR1 expression and CRH compared to those raised in the presence of adults or helpers (Taborsky et al., 2012). This was comlimented by a similar study by Taborsky in A. burtoni. Fry raised in social groups (i.e., naturalistic upbringing) had higher whole-brain GR mRNA expression compared to fry raised in pairs (Solomon-Lane and Hofmann, 2018). A follow-up study in N. pulcher examining specific regions of the brain showed that fry raised in the presence of adults and helpers had increased in GR1 mRNA expression in the telencephalon. This difference in results suggests that changes in GR1 expression are brain region specific. Antagonizing GRs in fry reared without adults and helpers resulted in more appropriate levels of aggression, indicating that the social effects of different rearing environments are mediated in part by GR (Nyman et al., 2017). Lastly, fry raised in the absence of adults and helpers displayed more neophobia in behavioral tests, which is indicative of higher stress in new environments (Bannier et al., 2017).

Table 1.2: Summary of Taborsky's work comparing *N. pulcher* fry raised in the presence and absence of adults and adolescents.

Phenotype	Fry Raised with	Fry Raised without
	Older Fish	Older Fish
Social competency	High	Low
Circulating CRH	Low	High
Whole brain GR1 expression	Low	High
Telencephalic GR1 expression	High	Low
Anxiety-like behavior	Low	High

$1.2.5 \quad A statotila pia \ burtoni$

A. burtoni, also referred to as Haplochromis burtoni, is a highly social circhlid species known for its extremely plastic phenotypes related to dominace. There exists two distint dominance phenotypes in males: those with territory that are reproductively active and those without territory that are reproductively suppressed (Fernald, 1977). Males with territory are dominant and have bright blue and yellow body coloration with a thick black stripe that passes over their eyes. They have vertical black stripes along their body, a black spot on their gill cover, and a red splotch just caudal of that. Non-territorial males are subordinant and resemble females, with camouflage coloration. The difference in dominance is also correlated with a difference in stress and hormonal regulation (Renn et al., 2008). Dominant males have increased testosterone and gonad size, while subordinant males have increased levels of cortisol

and experience faster growth. Males are able two switch between these phenotypes throughout their lives in response to changes in their social interactions.

Female A. burtoni also exhibit social dominance, though unlike the males, both dominant and subordinant females are reproductively active. Females take part in a brood care strategy known as mouthbrooding. The female lays eggs and immediately picks them up, storing them in her buccal cavity. Shortly after this, the males fertilizes the eggs. The mother keeps the offspring in her mouth for about two weeks as the eggs develop into fry. After the initial release, fry remain close to their mother and return to their mother's mouth in the presence of danger. Brooding mothers voluntarily starve themselves; however, in stressful situations they will eat their offspring. Mouthbrooding helps to protect the developing fry from predators and conspecifics, potentially reducing their exposure to stressful expreiences.

1.2.6 Current Investigation

The work of Barbara Taborsky has demonstrated that social fish species are plastic to their early life experience, showing changes in behavior and stress hormone expression as a result of social environment. Taborsky's work, however, is focused on the effect that social environment has on social behavior. There is little known about how maternal separation affects mouthbrooding fish. It is unknown to what extent mouthbrooding influences the neurophysiological development of fry. By comparing the effects of this child-rearing strategy to what is known in mammals, we can better understand the evolution of neuronal plasticity in response to early life experience, especially as it relates to parental care.

A recent thesis by Destine Krenik examined the role that maternal separation in A. burtoni had on behavior and neural GR expression (Krenik, 2018). Offspring were either reared with their mothers or were separated as eggs and raised without any adults. The thesis examined anxiety-like behavior and aggression in both conditions, but did not find any significant treatment-based differences. The brains of the fish were extracted and divided into forebrain, midbrain, and hindbrain for GR mRNA analysis. Krenik found significantly lower GR mRNA expression in the hindbrain of fry separated from mothers. The following research is conceptually based on this thesis with three major methodological changes. The first is that the paradigms used to measure social and anxiety-like behavior are different. The second change is that this thesis will examine whole-brain GR mRNA expression rather than section the brain into three regions. Lastly, the current research uses three different GR primers,

as opposed to Krenik's use of a single primer, in order to quantify more splice variants. The rationale for these changes is to collect different data for the same overarching question in order to build a better understanding as to how maternal separation alters GRs.

Chapter 2

The Experiment

2.1 Methods

2.1.1 The Fish

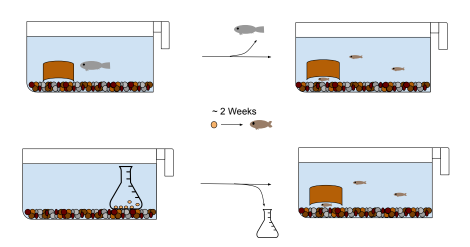


Figure 2.1: Experimental conditions.

Mouthbrooding fish were selected to be in either of two experiemntal groups. In the first condition (top) mothers were transfered to individual tanks containing gravel and a terra cotta piece. After about two weeks when fry were able to leave their mothers' mouths, the mothers were moved back into their home aquaria. In the second condition (bottom), eggs were removed from the buccal cavity and placed in a flask within an individual tank. After about two weeks the fry were able to freely swim and the flask was removed.

The parental generation of the focal juveniles originated from a wild-caught stock of *A. burtoni* collected from Lake Tanganyika in East Africa. Social groups containing males and females of the same generation were kept in aquaria at a temperature of

28°C and a pH of 8. Each aquarium's bottom was covered in gravel, and terra cotta pot pieces were placed in the tank to act as shelters and territory markers. Once a female fish began mouthbrooding, she was removed from the aquarium and randomly placed into one of two experimental conditions.

In the unseparated condition, mothers were removed from their home tank, weighed, and measured. They were then placed in small tanks containing gravel and a piece of terra cotta pot. Mothers continued to brood their young until the fry were old enough to regularly leave their mother's mouth (2 weeks post-fertilization), at which point the mother was removed from the tank to prevent her from eating the fry.

In the separated condition, mothers were weighed and measured and then the eggs were removed from their mouths by gently pulling down the bottom lips. The eggs were then placed in a 250 mL flask within a tank containing gravel. Once the eggs developed into freely moving fry (2 weeks post-fertilization), the flask was removed from the tank and a piece of terra cotta was added.

2.1.2 Behavioral Tests

Behavioral testing began approximately 130 days after the brooding mothers were placed into experimental conditions. Both assays were performed on the same day. Directly after each fish took part in the boldness assay, the brood was observed in the social assay. Prior to the start of behavioral testing, the focal brood was moved in its home cage to the testing area. The fish were allowed to adjust to the change in lighting for 10 minutes. Experimental tanks had the back and sides covered with white paper to minimize external visual stimuli. A video camera was placed approximately two feet from the front side. No humans were present in the room during behavioral recordings.

Boldness Assay

Boldness, or willingness to explore novel and open environments, is often used as a measure of stress. Animals that are stressed tend to freeze in place, seek cover, and avoid open spaces. Focal fish were individually placed in a novel aquarium containing gravel and a terra cotta shelter. A piece of red tape was horizontally placed on the outside of the tank, dividing it into top and bottom. The fish were allowed to acclimate for 10 minutes in the experimental tank before their behavior was scored for another 10 minutes. Boldness was measured on three axes: time spent in top half of tank, time spent in the bottom half of the tank, and time spent under the shelter. The fish was considered to be in a given region once its eyes crossed the border.

2.1. Methods

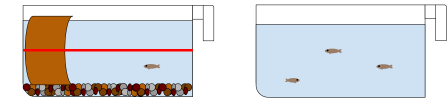


Figure 2.2: Behavioral paradigms.

Two paradigms were used to assess behavioral phenotypes of the fish. In the boldness assay (left) individual fish were placed in a novel tank containing gravel and a terra cotta shelter. After 10 minutes of habituation, time spent in the top half of the tank, in the bottom half of the tank, and behind the shelter was recorded for an additional 10 minutes. In the aggression assay (right) the whole brood was placed into a single, novel tank. As with the other assay, the fish were allowed to adjust to the change in environment for 10 minutes. The amount of chases, charges, and bites between fish were recorded for 10 minutes.

Aggression Assay

Because A. burtoni are highly social fish, it is relevent to assess how maternal separation affects their social behavior. Individual broods were transferred into a novel aquarium containing only water. The fish were allowed to acclimate to the new environment for 10 minutes before scoring began. The number of charges, bites, and chases between fish that occurred in 10 minutes were recorded. The total sum of aggressive behaviors were then divided by the number of fish in the brood to create a score. Broods that only had one fish in them were excluded from this paradigm.

2.1.3 Gene Expression Assay

Directly following behavioral testing, fish were measured for length and weight and quickly decapitated. The brains were then extracted and placed into 1 mL of RNALater and stored at 4 °C until needed for RNA isolation. RNA from each individual's whole brains was extracted using a Maxwell 16 LEV simplyRNA Tissue kit. RNA quality and concentration were confirmed using gel electrophoresis and nanodrop. For each sample, 100 ng of isolated RNA was reverse transcribed into cDNA with an Invitrogen Reverse Transcription kit.

Previously validated primers for GR1a, GR1b, and GR2 were used for qPCR. The housekeeping gene rpl32, which is a ribosomal protein coding gene, was used as a reference for GR expression. Each reaction contained cDNA (1 μ L), 2x Immuno

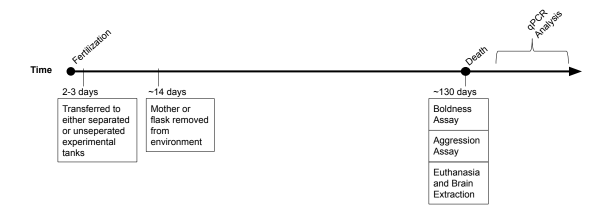


Figure 2.3: Timeline of experiment.

Two to three days after fertilization of eggs occurred, the broods were placed into one of two conditions: maternal separation or no maternal separation. After approximately two weeks, the fry were old enough to freely swim around their tanks. At this point, depending on the condition, either the mother or the flask was removed from the tank. The fry were allowed to age for about 130 days, at which point they were exposed to the boldness and aggression assays. Immediately after the behavioral testing, the fish were decapitated and their brains were extracted. RNA extraction and qPCR analysis took place after a variable amount of time.

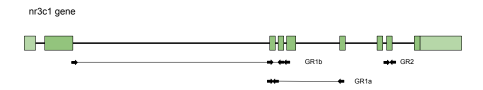


Figure 2.4: Primer design.

Primers sequences were taken from a study by . The location of the primers were then found by comparing the sequences to A. burtoni transcriptional sequences using the NCBI BLAST database. The forward primer of GR1a spans the

2.1. Methods

Mix (10 μ L), forward and reverse primers (1 μ L of 100 nM each), 50x SYBR Green (0.5 μ L), and nuclease-free water (6.5 μ L). The program used for the qPCR reaction involved a hot start (95°C, 5:00 min), 40 cycles that each ended with a fluorescent plate read (94°C, 0:45 min; 60.9°C, 1:00 min; 72°C, 0:30 min), 72°C for 2:00 min, and a melt curve analysis from 65°C to 95°C that measured fluorescence at 0.5°C intervals every five seconds.

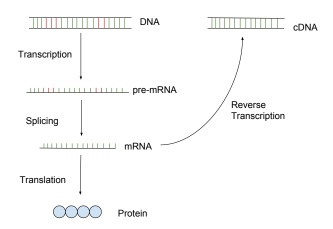


Figure 2.5: The central dogma of molecular biology including reverse transcription. The central dogma of molecular biology explains how DNA informs protein structure. DNA is transcribed into pre-mRNA. The introns (red), or non-coding regions, of the pre-mRNA are then spliced out, leaving only the exons (green). The spliced mRNA is then translated into a sequence of amino acids that together form a protein. Reverse transcription is the process of making mRNA into a double stranded nucleic acid known as cDNA or rtDNA. Unlike DNA, cDNA does not contain the sequences of introns.

Table 2.1: Genes and corresponding primer sequences used for qPCR

Target	Forward Primer (5' to 3')	Reverse Primer(5' to 3')
gr2	TGC CTC TGT CAC	AGT CGT CTG CGT
	TGC CAC CGT AG	CTG AAG TAA CTG
gr1a	TCA TAA GAT CTG	GTA GTT GTG CTG
	TTT GGT GTG CTC	GCC TTC AAC
gr1b	TGT TGG CTT CTC	GTT GTG CTG GCC
	CGG TTC ATC AC	ATC TGT GTT T
rfl32	TGC TGA TGC CCA	TCT TGG AGG AGA
	ACA TCG GTT	CAT TGT GGG

Troubleshooting

Prior to working with experimental fish, an age-matched brood was used to trou-

bleshoot behavioral testing. Following that, the brood was euthanized using MS-22 to practice brain extractions. Two of the extracted brains were used for gene expression troubleshooting. RNA was extracted from the brains and reverse transcribed into cDNA. All of the primer sets were tested using PCR on a gradient of melting temperature using this cDNA. The most effective melting temperature was then selected for qPCR. The efficiency of individual primers at the selected melting temperature was calculated with qPCR using a dilution series of the cDNA.

2.2 Results

- 2.2.1 Size
- 2.2.2 Boldness
- 2.2.3 Aggression
- 2.2.4 Glucocorticoid Receptor Expression

2.3 What Does this Mean?

2.4 Bibliographies

Of course you will need to cite things, and you will probably accumulate an armful of sources. This is why BibTeX was created. For more information about BibTeX and bibliographies, see our CUS site (web.reed.edu/cis/help/latex/index.html). There are three pages on this topic: bibtex (which talks about using BibTeX, at /latex/bibtex.html), bibtexstyles (about how to find and use the bibliography style that best suits your needs, at /latex/bibtexstyles.html) and bibman (which covers how to make and maintain a bibliography by hand, without BibTeX, at at /latex/bibman.html). The last page will not be useful unless you have only a few sources. There used to be APA stuff here, but we don't need it since I've fixed this with my apa-good natbib style file.

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