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1	Ontogeny and modulation, after PAMPs stimulation, of $\beta$ -defensin, hepcidin, and piscidin					
2	antimicrobial peptides in meagre (Argyrosomus regius)					
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4 5	Cindy Campoverde <sup>1, a</sup> , Douglas J Milne <sup>1, b</sup> , Alicia Estevez <sup>a</sup> , Neil Duncan <sup>a</sup> , Christopher J. Secombes <sup>b</sup> . Karl B. Andree <sup>a*</sup>					
6	<sup>a</sup> IRTA, Centro de San Carlos de la Rápita, San Carlos de la Rápita, 43540 Tarragona, Spain.					
7 8 9	<sup>b</sup> Scottish Fish Immunology Research Centre, University of Aberdeen, Zoology Building, Tillydrone Avenue, Aberdeen AB24 2TZ, Scotland, UK.					
10	<sup>1</sup> Contributed equally to the work.					
11	Contributed equally to the work.					
12 13	*Corresponding author: IRTA, Centro de San Carlos de la Rápita, San Carlos de la Rápita, 43540 Tarragona, Spain.					
14	Tel +34 977 745427, Fax 977 744138					
15	E-mail address: karl.andree@irta.es					
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#### Abstract

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Antimicrobial peptides (AMPs), components of innate immunity, play an important role in protecting fish. In this study we report the molecular cloning of full open reading frames and characterization of expression of three AMP genes (β-defensin (defb), hepcidin (hep2), piscidin (pisc) in meagre (Argyrosomus regius). A phylogenetic analysis of the expressed sequences obtained shows the defensin isoform forms a clade with the other members of the beta class of this family, hepcidin corresponds to hepcidin 2, and piscidin corresponds to class I of its respective family. Gene expression profiles of AMPs was investigated, by means of quantification of mRNA in nine development stages, from 8 days posthatching (dph) to accomplishment of juvenile form (120 dph). During development it was demonstrated defb, hep2, pisc were expressed in all stages of larval development and in juvenile tissues (kidney, spleen gut and gill). Moreover, expression patterns suggest the expression levels of theses AMPs are influenced by live prey (rotifer, Artemia) and first intake of commercial diet. Induction experiments in vivo (24 h) and in vitro (4, 12, 24 h) with PAMPs (LPS, poly (I:C), β-glucan) revealed significant changes in gene expression of the three AMP genes, in kidney, spleen, gut and gill. However, expression profiles differed in magnitude and time course response. defb expression shows a similar trend in vivo and in vitro in kidney at 24 h after LPS and β-glucan stimulation. The hep2 expression levels were up-regulated upon β-glucan challenge in vivo, more in gut and gills than kidney, while in vitro hep2 expression was upregulated in kidney cells by LPS, poly (I:C), β-glucan (4 h). pisc expression was up-regulated in kidney cells, splenocytes by β-glucan, but in gill cells by poly (I:C) and β-glucan in vivo. However, pisc expression was upregulated in kidney cells by β-glucan and gill cells by LPS at 4 post-stimulation in vitro. These data suggest that AMPs play an important role in defense against pathogens, with each AMP having differing efficacies against specific types of microorganisms, although follow-up studies focusing on the biological activities in fish are needed.

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- Keywords: antimicrobial peptides, PAMP, Argyrosomus regius, meagre, ontogeny, immune
- 53 response.

## Introduction

- Antimicrobial peptides (AMPs) are an evolutionarily conserved component of the innate immune response and recognized as a critical first-line defense against many pathogens. These innate effector molecules are present in both vertebrate and invertebrate life forms and play an important role in protection against a broad spectrum of pathogens including those of parasite, bacterial, fungal and viral aetiology [1] by molecular mechanisms of cellular disruption [2]. Fish is continuously fights against pathogens by secreting a wide range of AMPs [3] They are typically present in leukocytes (mast cells,
- 61 neutrophils), mucus cells (goblet and rodlet cells), cells lining epithelial surfaces and in gill, skin
- 62 intestine and other tissues [4–7].
- As a broad category of innate immune effector molecules, AMPs are divided into different families.
- 64 Several studies have found peptides in a wide variety of species belonging to the defensin, parasin,
- cathelicidin and hepcidin families, as well as piscidin, a family unique to teleost fish [8]. Many of the
- peptides identified have antibiotic activity as well as immunomodulatory functions. There has also been a
- 67 recent effort to study the potential of these peptides for applications in human medicine, and animal
- 68 welfare [3].
- 69 Fish AMPs are up-regulated in response to pathogen and appear to have direct broad-spectrum
- 70 antimicrobial activity towards both human and fish pathogens. However, the regulation the AMPs
- 71 transcripts in tissues in response to pathogen challenge is species and gene specific [9]. A major pathway
- of AMP production is via the recognition of pathogen associated molecular patterns (PAMPs) through the
- 73 pattern recognition receptors (PRRs), such as Toll-like receptors (TLRs), triggering an intracellular
- signaling cascade involving MyD88, TRAF6, IRAK1 and IKK. This signal promotes the activation and
- 75 translocation of NF-kB to the nucleus where it transcriptionally activates AMPs and other effector
- 76 molecules.
- 77 Defensins are a family of small cationic, amphipathic, cysteine-rich AMPs found in plants, fungi,
- 78 invertebrates, and vertebrates, including fish. These AMPs are involved in host-microbiota interactions,
- 79 immunomodulation, and provide strong links between innate and adaptive immunity. In vertebrates three
- 80 different Defensin subfamilies exist ( $\alpha$ -,  $\beta$  and  $\theta$ -defensins) that differ on the basis of the disulfide
- 81 bridges formed by the pairing and positioning of their six conserved cysteine residues [4,10]. However,  $\alpha$ -
- defensins are only found in mammals whilst  $\theta$ -defensins exist only in non-human primates [11]. In fish,
- 83 β-defensin paralogue sequences have been isolated in zebrafish (*Danio rerio*), fugu (*Takifugu rubriges*),
- 84 tetraodon (Tetraodon nigroviridis) and medaka (Oryziaas latipes) [12]. A recent study has also identified
- a novel group of anionic-defensins (fBDI-1 to -5) in flounder Paralichthys olivaceus that are expressed
- 86 during the larval period [10]. Constitutive β-defensin expression varies between species and depends on
- 87 the tissue type. Their biological functions have been investigated in relatively few fish species and have
- 88 broad-spectrum antimicrobial, antiviral and chemotactic ability, as reported for gilthead seabream (Sparus

aurata) and Atlantic cod (*Gadus morhua*) [13,14]. Remarkably, Defensins have also been implicated in some functions related to the reproductive system in orange-spotted grouper (*Epinephelus coioides*) [15].

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Hepcidins are another family of cysteine-rich peptides, structurally stabilised by disulphide bridges. They were first identified in human liver [16] and named liver-expressed AMPs (LEAP-1 and LEAP-2). The biological functions of mammalian Hepcidin include antimicrobial activity and regulation of iron metabolism. Hepcidin (Hep) has been extensively documented in many species of fish [8], where it has been shown to respond to bacterial infections and to function in antimicrobial activity. Whilst the highest expression level of hepcidin may be in the liver [17–19], as in mammals, in some fish species [20] these peptides are also expressed in other organs cardiac stomach, esophagus[17] heart, gill, spleen, kidney, peripheral blood leucocytes [18] dependent upon species. In mammals, two hepcidin isoforms are known. However, many teleosts (e.g., Perciform and Pleuronectiform fishes) possess up to seven isoforms of *hep*, as seen in winter flounder (*Pleuronectes americanus*) [17], rockbream (*Oplegnathus fasciatus*) [21] and redbanded seabream (*Pagrus auriga*) [22]. The presence of different environments and host-pathogen interactions [23].

Piscidins are AMPs that have cationic properties and an amphipathic α-helical structure. A total of 9 piscidins have now been identified in striped bass (Morone saxatilis) and 8 orthologous forms have been detected in hybrid striped bass [24]. They represent an evolutionarily conserved family of peptides unique to teleost fish and were first identified in the winter flounder with subsequent studies in hybrid striped bass (Morone saxatilis x M. chrysops) wherein the name was acquired [5,25]. They are widely active against Gram-positive and negative bacterial species, such as Streptococcus, Pseudomonas, Bacillus and Vibrio species [8], as well as being anti-fungal and anti-parasitic [26]. The primary mode of action is based on permeabilization of the plasma membrane of the pathogen. The immune-modulatory capacity of piscidins is another feature that has been widely assessed as they are able to modulate the expression of primarily pro-inflammatory genes as well as other immune-related genes, such as IL-1β, IL-10, TNF-α, NF-kb, NOS2, Myd88, TLR1, TLR3 and TLR4a through mechanisms involving a release of inhibition of NF-KB by down-regulation of IKB and, down-regulation of some anti-inflammatory signals such as IL-10, cAMP cascades and also indirect reduction of expression of inflammatory cytokines by elimination of pathogens [27,28]. Similarly, a study in orange-spotted grouper showed the immune-modulatory capacity of the epinecidin-1 (epi) gene by electrotransfer of the epi gene in the absence of bacterial infection, increases in the levels of transcripts encoding MYD88, TNF-a, TNF2, and NACHT [29-31]. Studies based on peptide structure, phylogenetic analysis, gene expression and antimicrobial activity have demonstrated that there are three different groups of Piscidin each of which may have different functions.

Meagre (*Argyrosomus regius*) is considered a major candidate species for large-scale fish farming in the Mediterranean and many regions of Europe, due to its favorable characteristics related to fast growth rate

and flesh quality. However, captive rearing of fish requires a well-developed larval rearing protocol and several factors affect the quality and survival rate of fish larvae, such as genetic background of the stock, diet, and the environmental conditions in which they are reared. This type of commercial growing necessitates intensive rearing in which crowding can augment skin abrasion/damage and lead to increased stress, facilitating transmission of infectious disease [32]. This study aims to elucidate one aspect of the innate response likely to be crucial in combating infectious bacterial diseases, namely some family members of AMPs that are present, and their level of expression in specific tissues. Target tissues were chosen for their significance as portals of entry of pathogens and their function in hematopoiesis, mucosal tissues of gills and intestine, and systemic tissues of kidney and spleen, respectively. We have characterized a *defb*, *hep* and *pisc* gene in meagre and describe their expression patterns during larval grow-out to evaluate when in development these peptides appear. In addition, we have examined how their expression is modulated by a variety of PAMPs, both *in vivo* and *in vitro*.

#### 2. Materials and methods

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- 137 2.1 Larvae production, and sample collection to determine AMP ontogeny
- Larvae used in the study were provided from meagre broodstock held in IRTA San Carlos de la Rapita,
- 139 Spain. The broodstock were maintained under controlled simulated natural water temperature and
- photoperiod using recirculation system (IRTAmar®). Breeders were fed frozen sardines (Clupeidae,
- fishmarket, Sant Carles de la Rapita, Tarragona, Spain), frozen Patagonian squid (Loligo gahi) (from
- Falkland Islands, Congelados Marcos, Tarragona, Spain) and a broodstock diet (Vitalis Repro, Skretting,
- Burgos, Spain). During the natural reproductive period (April-June), mature fish were selected based on
- oocyte size ( $< 550 \mu m$ ) and ease to extract sperm. Pairs of mature fish ( $21.2 \pm 3.7$  kg females and  $16.1 \pm$
- 2.6 kg males) were hormonally induced (15 µg/kg of des-Gly10, [D-Ala6]-gonadotropin-releasing
- hormone ethylamide, Sigma, Spain) to spawn spontaneously in 10,000 L tanks and spawned eggs were
- collected with a passive egg collector placed in the outflow of the tank. Egg number was estimated by
- 148 counting the total number of eggs in three sub-samples of 10 mL and fertilization by examining the
- development of 100 eggs. Batches of 50,000 eggs were incubated (18-19 °C) in 35 L mesh (300 μm
- mesh) bottomed incubators with aeration and gentle air-lift water exchange that were placed in 2,000 L
- tanks. Hatching rate was determined by estimating the number of larvae (three 100 mL sub-samples)
- obtained from the stocked eggs.
- Larvae were transferred and distributed into two 1.5 m<sup>3</sup> tanks under a mesocosm system. Water
- temperature was maintained at 20 °C. From two days post-hatching (dph), larvae were fed enriched
- rotifers until 8 dph. Freshly enriched Artemia metanauplii were introduced at 9 dph until 31 dph and a
- formulated artificial diet was incorporated at 21 dph until the end of the experiment (Fig. 1a). Samples of
- 157 larvae were collected from post hatch until juvenile for growth measurements. Random samples of larvae
- and juvenile were taken at 8, 15, 29, 40, 43, 60, 85, 96, and 120 dph for each sample point collected in an
- 159 Eppendorf tube on ice containing RNAlater™ (Ambion, Austin, Texas), then preserved at -80 °C until

- RNA extraction. Time points for analysis were chosen as proxies for the specific periods of change in rearing practices; primarily changes in diet, but also rapid changes in organogenesis, all of which increase stress and enhance susceptibly to diseases. The fish were euthanized using a high concentration of MS222 (1 g /L) (Aldrich, E10521) prior to sample collection. Larger larvae (post-29 dph) had excess tissue trimmed to reduce signal dilution from non-target tissues; the anterior section from the gills onwards, and
- the posterior part from the anus to the tail. The samples were collected from each time point for the study
- of AMP gene expression. At days 85, 96, and 120 individual tissues (gill, kidney, spleen, and intestine)
- were excised aseptically.
- 168 2.2 Total RNA extraction and reverse transcription
- Total RNA was extracted using Trizol reagent (Invitrogen) according to manufacturer's instruction. RNA
- 170 concentration and purity was determined by spectrophotometry (NanoDrop2000, Thermo Fisher
- Scientific, Madrid, Spain) measuring the absorbance at 260 and 280 nm. The quality of extracted RNA
- was verified with visualization of the 28S and 18S ribosomal RNA bands by agarose gel electrophoresis.
- 173 For preparation of cDNA total RNA was treated with DNase 1, Amplification Grade (AMPD1-1KT,
- 174 Sigma-Aldrich, Broendby, Denmark), according to manufacturer's instructions to remove possible
- 175 contaminating genomic DNA. Total RNA was reverse transcribed in a 20 µL reaction volume containing
- 176 2  $\mu g$  total RNA using the ThermoScript <sup>TM</sup> Reverse Transcriptase (Invitrogen) with oligo (dT)  $_{(12-18)}$  (0.5
- 177 μg/μL) and random hexamer primers (50 ng/μL) 10X RT buffer [200 mM Tris-HCl (pH8.4), 500 mM
- 178 KCl] 1.5 mM MgCl<sub>2</sub>, 800 mM dNTP mix, RNase inhibitor, SuperScript <sup>TM</sup> II RT, followed by RNAse H
- 179 (Invitrogen) treatment. Reverse transcription reactions were prepared, which were placed in a
- thermocycler (Mastercycle ® nexus GSX1, Eppendorf AG, Hamburg, Germany) and run according to
- manufacturer's instruction. Negative controls (no RT enzyme) were included to confirm absence of
- genomic DNA contamination. The samples were then diluted 1:20 in molecular biology grade water and
- 183 stored at -20 °C.
- 184 2.3 Gene Isolation
- Meagre were sampled for collecting data on specific growth rate of the chronological samples. The
- samples were collected in RNAlater at each time point, for extraction of RNA to be used in gene
- 187 expression analysis. Sequences of target genes from the online database Genbank were chosen from
- extant marine teleost species that were available, such as large yellow croaker (*Larimichthys crocea*),
- mandarin fish (Synchiropus splendidus), Humphead snapper (Lutjanus sanguineus), European seabass
- 190 (Dicentrarchus labrax), orange spotted grouper, but also tilapia (Oreochromis spp). The sequences were
- 191 aligned, using CLUSTAL W incorporated in the package BioEdit
- 192 (http://www.mbio.ncsu.edu/BioEdit/bioedit.html), for designing degenerate/consensus primers for
- amplification from cDNA of tissues. The fragments amplified were separated by gel electrophoresis and
- resulting bands of the expected length were excised, purified (QIAQuick PCR purification kit, Qiagen)

- and sequenced (Sistemas Genomicos, Valencia, Spain). The transcript sequences thus obtained were used
- as templates to design primers for specific q-PCR assays.
- 197 For some target genes, there are suspected to be paralogs that could create "noise" for the gene expression
- analysis if the gene specific primers targeted only the most conserved regions. To avoid this 5′-3′ RACE
- reactions were performed to obtain the full open reading frame using the method described by Zou et al.
- 200 [33]. The RACE reactions followed the initial fragment isolation using degenerate-primed PCR
- 201 with the primers: Pisc F TGGTTGTTCTCATGGCTGAAC, Pisc R GGTCA
- 202 TAAGAAAGTGAACGT, Hep F ATGAAGACATTCAGTGTTGC, Hep R-
- 203 CAGCAACCGCAGCAAA, Defb F GTGCTTCTCCTGATGCTCGC, and Defb R -
- 204 CTGTATCTTCGAGGGCAAC.
- 205 2.4 Phylogenetic Analysis
- 206 Confirmation of the relation of the AMP sequences obtained to extant marine teleost sequences was
- performed using a phylogenetic analysis. Evolutionary analyses were conducted in MEGA5 [34] utilizing
- only the coding portion of the open reading frames of each AMP. The nucleotide substitution model used
- was chosen utilizing the tool incorporated in MEGA5. The evolutionary history was inferred by using the
- 210 Maximum Likelihood method based on the Jukes-Cantor model [35]. Initial tree(s) for the heuristic
- search were obtained automatically as follows. When the number of common sites was < 100 or less than
- one fourth of the total number of sites, the maximum parsimony method was used; otherwise, the BIONJ
- 213 method with MCL distance matrix was used. The analysis involved 33 nucleotide sequences. All codon
- 214 positions were included in the analysis, however, positions containing gaps and missing data were
- eliminated. There were a total of 114 positions in the final dataset.
- 216 2.5 PAM stimulation
- To investigate the effect of PAMP stimulation on the expression of *defb*, *hep* and *pisc*, 38 healthy juvenile
- 218 individuals (30 40 g each) of meagre were held in a recirculating water tank (100 L) with seawater at 20
- °C. The fish were fed with a commercial diet twice a day (9:00 h and 16: 00 h) for two weeks to acclimate
- 220 them to the indoor culture environment. For primary cell culture and collection of samples, fish were
- killed after anesthesia with 50 mg/L MS-222.
- 222 2.5.1 AMPs expression after in vivo stimulation
- Fish were injected intraperitoneally with 100 uL PBS containing 100 µg poly (I: C) (3.3 mg/kg) (Sigma,
- UK. P1530), 400 μg LPS (13.3 mg/kg) (Sigma, UK. L3129) or 100 μg β-glucan (3.3 mg/kg) (Sigma, UK.
- 89862). The control animals were injected with PBS only. After 24 h, the individuals were dissected and
- 226 tissues (kidney, spleen, gut and gill) sampled. Total RNA was isolated and cDNA was prepared as
- described above. The mRNA expression level of *defb*, *pisc* and *hep* were determined by q-PCR. Relative
- 228 expression was normalized to GAPDH expression and calculated as arbitrary units and converted to a
- proportion relative to the PBS control samples.

- 230 2.5.2 AMPs expression after in vitro stimulation
- Tissues (kidney, spleen, gill, intestine) from apparently healthy fish were passed through a 100 µm nylon
- mesh cell strainer (SefarNytal PA-13xxx/100, Spain) in Leibovitz L15 medium (Gibco) containing
- 233 penicillin/streptomycin (Gibco, #15140-122) at 1:1,000 and 2% foetal calf serum (Gibco, #10270-098).
- The resulting cell suspension was collected and centrifuged at 400 x g for 10 min. The supernatant was
- removed and replaced with 10 mL of the previously described L15 media. The cell suspension was again
- centrifuged and supernatants removed and replaced with 30 mL of media. Cells were distributed to 12
- well microtiter plates in 5 mL aliquots. Wells were stimulated using LPS (Sigma, #L3129-100MG) at 50
- 238  $\mu g/mL$ , poly (I:C) (Sigma, #P1530-25MG) at 100  $\mu g/mL$ , and  $\beta$ -glucan (Sigma, #89862-1G-F) at 50
- 239 μg/mL in triplicate. Control samples included 250 μL of PBS. Four, 12 and 24 h after stimulation the
- 240 cells were harvested and centrifuged at 400 x g for 10 min., the supernatant discarded, and the pellet
- suspended in RNAlater. Total RNA was isolated and cDNA was prepared as described above.
- The q-PCR reactions for AMP gene expression were carried out in duplicate on a LightCycler® 480 Real-
- Time (Applied Biosystems, Roche). A master mix was prepared from: 6 µL SYBR Green Supermix (Life
- Technologies), 10  $\mu$ M of each primer, diluted in molecular biology grade water (Sigma) and 4  $\mu$ L of
- 246 cDNA in a final volume of 10 μL. The real-time q-PCR cycling was carried out as follows: 10 min at
- 247 95°C, 40 cycles of 95 °C for 25 s, followed by an annealing step of approximately 59 °C for 25 s
- 248 (annealing temperatures were adjusted for each specific primer pair), followed by 72 °C for 15 s, with a
- final melt curve stage of 0.5 °C increments from 75 °C to 95 °C. Each sample on the q-PCR plate had two
- 250 technical replicates. Primer sequences used and amplicon lengths for each assay are shown (Table 1). The
- specificity of the primers was checked by running a q-PCR and confirming that only one melt peak was
- produced and also by running a subsample on an agarose gel (2%) to confirm the presence of a single
- band of the expected size. The efficiency of amplification (E %) of each primer pair was assessed from
- five serial ten-fold dilutions of cDNA from individual tissues, then calculated following the equation: E%
- $= 10^{(-1/\text{slope})} 1$  where the "slope" is that calculated from the regression line of the standard curve.
- Efficiencies of the gene expression assays ranged from 101.95% to 104.3% ( $102.85 \pm 1.27$ ) (Table 1).
- 257 The absence of primer dimer formation in the NTC was also confirmed. Relative expression of genes
- from larvae, and each tissue from juveniles (ontogeny study), was normalized using the three endogenous
- 259 controls glyceraldehyde phosphate dehydrogenase (GAPDH), beta-actin (β-actin), and hypoxanthine-
- 260 guanine phosphoribosyltransferase (HPRT) as determined using geNORM
- 261 (http://www.primerdesign.co.uk/products/9461-genormplus-kits/), while the relative quantification at 120
- dph, showing the least expression, was used as the calibrator.
- For analyzing the relative expression in vivo and in vitro after PAMP stimulation, data was normalized to
- 264 GAPDH expression, as this endogenous control gene showed less variability under the conditions tested.

- After normalization to this endogenous control as arbitrary units, results were converted to a proportion
- relative to the control group (PBS injected fish).
- 267 2.7 Statistical analysis
- All data were checked for homogeneity of variances using a Levene's test by univariate analysis in a
- general linear model, based on a Tukey HSD post-hoc test, with a sample size of n = 10 (larva) n = 8 (
- juvenile) to determine differences between time points ( $P \le 0.05$ ). The Kruskal-Wallis test was used for
- analyzing the expression for the *in vivo* (n = 8) and *in vitro* (n = 6) samples after PAMP stimulation, using
- the statistical software package SPSS 20.0 (SPSS Inc., US).
- **273 3. Results**
- 274 3.1 Gene Isolation
- 275 Specific gene products were obtained using degenerate or consensus-primed PCR assays. All sequences
- were analyzed using the BLAST utility via the National Center for Biotechnology Information (NCBI)
- database (http://www.ncbi.nlm.nig.gov/) to confirm their identity as proper orthologs of the intended
- 278 target gene. Sequencing followed by BLAST analysis provided presumptive identification of AMP
- transcripts for *defb*, *hep* and *pisc*, reported for the first time in this species. Sequences were also obtained
- 280 for GAPDH, β-actin and HPRT as endogenous control genes for gene expression assays for which
- 281 sequences have been provided in GenBank under the following accession numbers MF186586,
- MF186587 and MF186588, respectively. The AMP sequences have been uploaded to GenBank under the
- following accession numbers and length the fragment defb = MF074072 (195 bp), hep2 = MF074074
- 284 (255 bp) and pisc = MF074073 (216 bp).
- 285 3.2 Phylogenetic analysis
- To verify the relationship of meagre AMPs to known AMP genes, the A. regius sequences were used to
- construct a phylogenetic tree with extant marine teleost sequences of AMPs (when available) (Fig. 2).
- 288 Results indicate the meagre hepcidin gene is more similar to that of hep 2 from Pseudosciaena crocea and
- species of the moronidae family (*Dicentrarchus labrax* and *M. chrysops*). The *defb* sequences appear to
- be most closely related to mugilidae (*Liza haematocheila*). Meagre pisc was included in the class I clade
- of piscidins according to the scheme of Salger et al. [24], with affinity to Larimichthys crocea piscidin
- supported by a significant bootstrap value (94).
- 293 3.3 Gene expression analysis
- 3.3.1 Ontogeny of AMPs
- In the present study, AMP expression during larval development has been assessed. Meagre growth was
- described by the equation LS =  $0.0016 \text{ dph} + 0.7219 \text{ dph} 3.9013 (R^2 = 0.9851)$  and wet weight by the
- equation W.Wt = 1.7368 dph 66.966 dph + 317.74 ( $R^2 = 0.9654$ ) between 0 and 133 dph of the
- 298 experimental rearing (Fig. 1b).

- The expression of *defb* exhibited marked differences between the larval and juvenile stage, with a statistically significant increase in expression during transition from *Artemia* to standard commercial dry pellet feed at 29 dph, followed by a gradual decrease until 60 dph. Expression patterns of *defb* transcripts in individual tissues show significantly different expression profiles from 85 dph to 120 dph. The highest expression was seen in kidney compared to other tissues tested, however, in spleen there was no difference in expression over time. (Fig. 3a).
- The *hep2* expression in larvae remained markedly low from day 8 to day 15, where the expression was increase at day 29, when the switch from feeding on *Artemia* to artificial feed is nearly complete. In the analysis of separate immune tissues a gradual increase of expression was observed over time (85, 96, or 120 dph) in spleen, gill and gut. Although the trend of increase in expression over time is similar in all three of these tissues, a difference in mRNA level was only statistically significant in gut, where an increase was seen at 96 to 120 dph. However, *hep2* gene expression was highest in kidney compared with the other tissues (gill, gut, spleen) (Fig. 3b).
- The *pisc* transcripts showed a pattern of expression markedly different across developmental stages. The *pisc* gene expression remained low from 8 to day 29, where the expression increased at day 40, when the larvae feed includes only artificial feed, followed by a gradual decrease thereafter. In isolated tissues, high *pisc* expression was detected in gills, but similar patterns of expression were found in kidney and spleen at 85, 96, 120 dph. However, gut tissue had consistently low levels of *pisc* expression during the time points examined (Fig. 3c).
- 3.3.2 Response of meagre AMPs genes to **in vivo** PAMP stimulation
- After injection of fish with PAMPs, the expression levels of *defb*, *hep2* and *pisc* were examined by qPCR at 24 h post-stimulation. The *defb* transcription was up-regulated in kidney post-injection with LPS and  $\beta$ -glucan while a significant down-regulation was seen in spleen, gut and gills following LPS injection, and in spleen following poly (I:C) injection (Fig. 4a). In contrast, *hep2* was significantly up-regulated in gut and gills by  $\beta$ -glucan, and in gills by LPS (Fig. 4b). Lastly, the transcription of *pisc* showed significant up-regulation in kidney, spleen and gills by  $\beta$ -glucan treatment, and there was also significant up-regulation in gills by poly (I:C) stimulation (Fig. 4c).
- 3.4.3 Response of meagre AMPs genes to in vitro PAMP stimulation
- After stimulation of isolated leucocytes in vitro with PAMPs, the kinetics of changes in expression levels of *defb*, *hep2* and *pisc* were examined by qPCR. No change in expression of *defb* gene was found in the stimulated kidney cells at 4 h and 12 h, but at 24 h there was significant up-regulation ( $P \le 0.05$ ) using poly (I:C), LPS and β-glucan, greater than 20-fold in the latter case. With splenocytes no changes were seen at 4 h or 24 h, but at 12 h a small decline in the level was detected after stimulation with poly (I:C), LPS and β-glucan. The mRNA expression of *defb* in gill cells was significantly up-regulated at 4 h after stimulation by poly (I:C), but thereafter it showed a decline at 12 h 24 h. Similarly, *defb* was up-

- regulated with LPS stimulation at 12 h, but declined at 24 h. β-glucan stimulation resulted in down-
- regulation from 12 h onwards. Lastly, the mRNA expression of *defb* in intestinal cells was significantly
- up-regulated ( $P \le 0.05$ ) at 4 h and 12 h post-stimulation with poly (I:C), LPS and β-glucan, however an
- up-regulation at 24 h was only seen with LPS and β-glucan stimulation (Fig. 5a).
- For hep2, transcription in kidney cells was significantly up-regulated at 4 h post-stimulation with poly
- 339 (I:C), LPS and β-glucan and, thereafter, it showed a drastic decline at 24 h in the presence of these three
- PAMPs. In splenocytes only at 24 h after stimulation by PAMPs was a significant effect seen, with down-
- regulation apparent for all three PAMPs. In gill cells, the gene transcripts were rapidly and significantly
- up-regulated at 4 h after stimulation by poly (I:C), but showed a decline at 12 h, and later returned to
- basal levels at 24 h. In intestinal cells, the gene transcripts were significantly up-regulated at 4 h after LPS
- and β-glucan treatment, had declined at 12 h, but were again enhanced at 24 h after LPS stimulation (Fig.
- 345 5b).

- A significant up-regulation of *pisc* expression was seen in kidney at 4 h and 12 h after stimulation by  $\beta$ -
- glucan and at 24 h after stimulation using poly (I:C). In splenocytes, up-regulated was seen at 24 h after
- 348 stimulation by LPS. The gills cells displayed a higher expression level at 4 h and 12 h after stimulation
- with LPS, and at 4 h with  $\beta$ -glucan, compared with the control. Significant down-regulation of the
- expression of *pisc* transcripts was observed in the intestine, but only at 12 h post-stimulation with the
- 351 PAMPs (Fig. 5c).

# 4. Discussion

- 353 Antimicrobial peptides are a diverse group of small peptides that exhibit broad-spectrum antimicrobial
- activity towards pathogens (parasites, bacteria, viruses and fungi). They function as a part of the primary
- defense within the innate immune system protecting the host from attack by foreign particles and
- organisms in a non-specific manner [36]. Fish are free-living organisms from the initial stages of
- development and exposed to numerous pathogens before adaptive immunity has fully developed. It is
- known that some aspects of teleost immunological capacity are limited in early development and they rely
- on innate mechanisms such as AMPs for survival. This is the first description of the temporal appearance
- and flux of expression during larval development of representative members of three AMP families of
- immune effectors from A. regius, including Defensin, Hepcidin, and Piscidin. A phylogenetic analysis of
- the expressed sequences obtained (Fig. 2) shows that among the transcripts from A. regius, the defensin
- isoform forms a clade with the other members of the beta class of this family, hepcidin corresponds to
- 364 *hepcidin* 2, and *pisc* corresponds to class I of its respective family.
- 365 The phylogenetic tree reveals that the isoform of *def* identified from meagre is *defb*, although no homolog
- from L. crocea was found in the GenBank database so the formation of a clade with defb from Liza
- 367 *haematocheila* is likely an artifact of the database content. During growout the ontogeny of the expression
- of the identified AMPs from meagre was evaluated. In this study, *defb* was found expressed in all stages

of larval development and in juvenile tissues examined in meagre. In larva, expression levels were identified after complete absorption of the yolk sac and were found to be significantly up-regulated around the time of *Artemia* feeding though decreased after weaning. This increased expression after the start of the exogenous feeding may reflect a need for efficient defense mechanisms against potential pathogens that can enter with the live feed or simply up-regulation due to PAMP stimulation from commensal organisms in the feed. The *defb* expression at early developmental stages has been demonstrated in vertebrates and invertebrate [4]. Two β-defensin genes were identified in snout bream (*Megalobrama amblycephala*) during early development of embryos and fry [37]. Investigation of the multiple DefB isoforms were found in olive flounder (*Paralichthys olivaceus*) after hatching, revealed a constitutive *FBD1* mRNA expression before and after hatching at 1-35 dph [10]. While only one isoform was identified in the current study, it also exhibited constitutive expression as seen with *FBD1* mRNA. In common sole (*Solea solea*) *defb* transcripts showed high transcription level across all larval stages [38]. In this context, expression patterns observed in this study might suggest that this peptide is implicated in innate conditioning of the early larval stages, preventing pathogenic assault, and conditioning of the gut microbiota [39].

 The  $\beta$ -defensin gene expression in teleosts has been assessed in tissues from healthy fish, but the levels and patterns of expression varied for individual defensin genes [13,37]. In the present study with meagre, analysis of the isoform obtained revealed expression remained somewhat constitutive, but was highly expressed in kidney with lower levels in gill, gut, and spleen at 120 dph. The decrease in expression seen between 29 dph and 60 dph may be due to the expression signal being diluted as the larvae grew, as the abundance of *def*-positive cells likely did not increase in the same proportion as the general growth and increase in musculature. However, the isoform under study can influence the expression pattern as well as the host species. In addition to viable microbial pathogens, the components of a fish's diet can act to regulate *defb* expression. In common carp, *BD1* and *BD2* in gills and *BD1* in skin were found significantly higher upon addition of  $\beta$ -glucan to the food [40]. As studies with Gilthead seabream fed microalgae (*Nannochloropsis gaditana*, and *Tetraselmis chuii*) show, an increase in the expression level of *defb* in head-kidney and gut can be accomplished via dietary supplement [41]. On the basis of the results obtained in this study (Fig. 3), further dietary studies to investigate how change of food from live prey (*Artemia*) to commercial feed may influence AMPs expression in larvae.

The potential to modulate the expression level of the AMPs (*defb*, *hep 2*, *pisc*) was also analyzed after *in vivo* and *in vitro* immune activation by PAMPs (LPS, poly (I:C), β-glucan). The baseline expression for all three types of AMPs was lowest in isolated gut tissue. This has biological sense when considering the importance of maintaining a healthy gut microbiota. A leaky promoter or slight defects in the secretion systems could have disastrous consequences for the health of the host [42]. Maintaining a tight regulation of control would help to avoid incurring inadvertent damage to the gut microbiota, therefore the control in the gut is more likely to be bi-model rather than multi-model [42]. Our results showed that the expression

of *defb* was significantly up-regulated in the kidney after injection with LPS and  $\beta$ -glucan at 24 h. This result is in accordance with the idea that the kidney is an important hematopoietic organ and filters the blood where strong up-regulation of *defb* can be expected as a prophylactic measure to ensure pathogen elimination. Similar results have been seen in Atlantic cod, where *defb* is up-regulated in the kidney 48 h post-challenge with *Vibrio anguillarum* [43], and in the kidney in olive flounder 1 h after infection with *Edwardsiella tarda* [10]. Using cells isolated from individual tissues our results in meagre also show up-regulation of AMPs following PAMPs stimulation. The expression of *defb* was increased in kidney and gut cells at 4 and 24 h after stimulation by  $\beta$ -glucan, LPS and poly (I:C). These results suggest early responses (4-12 h) of *defb* in meagre are more dependent on the stimulus in mucosal tissue than in systemic tissues where significant differences were not observed. Although, later response (24 h) in the kidney were more pronounced. Interestingly, similar trends were found *in vivo* and *in vitro* with regard to *defb* expression in kidney at 24 h after LPS and  $\beta$ -glucan stimulation, possibly reflecting a large population of antigen presenting cells responsive to PAMPs in this tissue.

The phylogenetic analysis also reveals that meagre hep2 and pisc were more similar to equivalent molecules in Larimichthys crocea as expected, since this is taxonomically a quite similar species of fish. Hepcidin is highly conserved from fish to humans [44,45]. In fish, it plays a major role in the innate immune response and iron regulation. In the present findings, meagre hep2 gene expression patterns during larval development showed high expression levels between 8 dph and 15 dph, then increased significantly at 29 dph at the end of metamorphosis. Similar results were observed in Solea solea where SsoHAMP expression levels increased from 8 dph [38], as well as in Atlantic cod [46] and half -smooth tongue sole [47]. However, in the redbanded seabream and in Pseudopleuronectes americanus hep isoforms were found to peak earlier between 5 and 15 dph [17,22], whilst in Medium carp (Puntius sarana) a hep homologue had stable expression from 6 h post-fertilization onwards [48]. During these timings, numerous changes occur in embryogenesis and early larval development. Elevated expression at this time likely indicates a preparation of the larvae for dealing with a pathogen-laden environment after hatching, as well as conditioning the gut mucosa by stabilization of the normal commensal microbiota that prevent colonization of the gut by opportunistic pathogens through spatial exclusion [39]. Physical barriers of the innate immune response are not sufficient by themselves and when these barriers should be insufficient, additional mechanisms, such as the production of AMPs, are required. From the moment of hatching the fish larvae needs adequate defense mechanisms, with the commencement of external feeding potentially exposing larvae to viable biological agents contaminating the feed.

In this study, *hep2* was expressed in all tissues examined, and highly abundant in kidney followed by spleen and gills. Earlier studies from other species, have shown that *hep* transcripts are usually observed at a higher level in the liver in fish, although they can also be detected in other tissues [43-46]. Indeed, the expression patterns between different fish *hep* is highly divergent and may be related to the specific isoform(s) examined [18].

The in vivo challenge results indicated, that hep2 was also found to be up-regulated upon  $\beta$ -glucan stimulation, but in gut and gills rather than kidney. Further, in gills hep2 was significantly up-regulated by LPS stimulation. Studies in fish species of Hep protein have shown multiple isoforms exist and these are classified into two groups: Hamp type 1 and Hamp type 2. One Hep isoform has iron regulatory roles sharing a high degree of homology with mammalian counterparts, and the second Hep copy has retained only the antimicrobial role [9,49]. Phylogenetic analysis in this study has shown the isoform identified in meagre is more similar to Hamp 2 and likely does not retain the iron regulatory functions. In this study the responsiveness of this isoform has only been demonstrated with the in vitro and in vivo PAMP stimulation indicating its antimicrobial functions. Additionally, there may exist other hep isoforms with different tissue-specific expression patterns related to distinct tissue-specific regulation via their promoters [9]. Indeed, different hep isoforms in a single species can have different levels of expression within the same tissue, and can be affected by different stimuli [21,22,50]. For example, in the olive flounder where multiple isoforms exist, Hep-JF1 was induced in a wide array of tissues (including liver, gill, kidney and spleen) by LPS, but another variant Hep-JF2 was unaffected [18]. However, whilst LPS is well known to modulate *hep* expression in teleost fish, in large yellow croaker (*Pseudosciaena crocea*) where spleen, heart, stomach, kidney, and liver tissues were examined, LPS stimulation significantly enhanced expression only in spleen, heart and stomach [20].

The in vitro challenge results indicated, hep2 expression was up-regulated in kidney and gut cells at 4 h post-stimulation by β-glucan and LPS, and at the same time in kidney after poly (I:C) treatment (4 h), with later (24 h) stimulation of expression in intestinal cells by LPS. However, the changes were relatively small as compared to what might be expected in hepatocytes which were not analyzed in this study. The hep expression has been shown to be up-regulated *in vitro* by several PAMPs in other species [51]. Also in cell culture, in rainbow trout, where hep mRNA level was increased in a macrophage (RTS11) cell line treated with poly (I:C) [52]. In tilapia, Hep2 and Hep3 (TH2-3) showed inhibition of proliferation and migration of a tumor cell line (HT1080 cells) in a concentration-dependent manner [53].

Class I Piscidin has been shown to be more effective against prokaryotes and ciliated protozoan infections [24], as compared to class II and class III. Although in this study functional analyses have not been evaluated, on the basis of q-PCR results, meagre class I Piscidin showed up-regulation with all three PAMPs both *in vivo* and *in vitro*, which is suggestive of it being more responsive to a broader range of stimuli. Piscidin is generally expressed early during fish development and increases in expression over time. Our results showed that *pisc* transcripts were detected in 8 dph larvae, with maximum levels seen at 40 dph when they have completed the transition to juveniles. There was a decrease after 40 dph, but as discussed above for *defb*, this decrease may in fact be related to a dilution effect due to rapid growth after the end of metamorphosis. In more developed juveniles where individual tissues were examined, *pisc* transcripts were detected at high levels in the gill, with lower expression in the kidney and spleen, and lowest expression in the gut. In studies of striped bass, Piscidin was observed by immunofluorescent

labelling in post-vitellogenic oocytes, embryos or larvae from 14 dph [54]. European seabass Piscidin 3 was detected in cytoplasmic granules of inflammatory cells, and mast cells [55]. Until recently, piscidins had only been found in fish species from the superorder Acanthopterygii, but there has recently been *pisc* genes identified in Atlantic cod [56], where expression is seen in more diverse tissues including liver, blood, gall bladder, pyloric caeca, stomach, rectum, muscle, heart, and brain [56–58], as well as in mast cells and rodlet cells of the skin, gill and intestine [59].

In the present study *in vivo* stimulation also modulated *pisc* expression, with the highest level of expression found in kidney, spleen and gills at 24 h after injection with  $\beta$ -glucan, and in gills after poly (I:C) treatment. In other fish species *pisc* genes can be induced by a variety of stimuli such as LPS [60,61], parasite antigens, poly (I:C) [62] and  $\beta$ -glucan as a representative PAMP from fungi [58]. For example, in Mandarin fish (*Siniperca chuatsi*) a significant up-regulation of *pisc* was seen in most tissues, except the liver, after stimulation with LPS [63], while in large yellow croaker, *pisc* expression was up-regulated in response to the ectoparasite, *Cryptocaryon irritans* [26]. Furthermore, in hybrid tilapia (*Oreochromis spp.*) Piscidin showed a correlate with lower mortality after injection with *V. vulnificus* [64]. Regarding *pisc* expression *in vitro*, this was up-regulated in kidney cells by  $\beta$ -glucan and in gill cells by LPS at 4 and 12 h post-stimulation. A late induction in spleen and kidney cells was also seen in response to LPS and poly (I:C) at 24 h post-stimulation. Interestingly, *pisc* expression *in vitro* (4-12 h) and *in vivo* (24 h) showed up-regulation in the same tissues from whole animals and isolated cells (kidney and gills) by  $\beta$ -glucan stimulation. These results demonstrate that timing of induction from each stimulant has tissue-specific differences, although the *pisc* isoform identified can be induced by all three types of PAMPs.

In conclusion, the current study is the first identification of AMPs from meagre and demonstrates their expression modulation during development and in response to PAMPs. Further studies will be needed to elucidate whether additional isoforms exist and the antimicrobial activities of the peptide proteins. The *A. regius defb, hep2*, and *pisc* transcripts were expressed at different levels in early developmental stages and tissues of juvenile meagre, and the data hint that expression is regulated or influenced by dietary factors such as switching to live feed, or weaning onto artificial diets. This may allow larval feed composition for *A. regius* larvae to be manipulated to augment AMP gene expression to improve antimicrobial defenses as has been seen for *defb* expression in common carp [40] and gilthead seabream [41]. This work provides useful information for future investigations of the innate defense mechanisms in early development of meagre and studies focusing on the biological activities of these antimicrobial peptides for this fish species.

## Acknowledgements

511	CC received funding from a Ph.D. grant awarded by the Secretaría Nacional de Educación Superior
512	Ciencia, Tecnología e Innovación (SENESCYT), Ecuador. This project has received funding from the
513	European Union's Seventh Framework Programme for research, technological development and
514	demonstration (KBBE-2013-07 single stage, GA 603121, DIVERSIFY). We would like to thank G
515	Macià, M. Monllaó, and S. Molas for their help in live prey production.

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**Table. 1** – Primers used for gene expression analysis by real-time q-PCR, including the amplicon size and primer sequences. Abbreviations: gapdh; glyceraldehyde 3-phosphate dehydrogenase; β-actin, beta-actin; hprt, hypoxantine-guanine phosphoribosyltransferase; β-def, β-defensin; hep, hepcidin; pisc, piscidin.

	Genes	E (%)	Size (bp)	Forward primer (5'->3')	Reverse primer (5′->3′)
	gapdh	100.00	109	CCAGTACGTGGAGTCCACTG	AGCGTCAGCGGTGGGTGCAGAG
Reference	β-actin	100.05	212	TGGGGGAGCAATGATCTTGATCTTCA	AGCCCTCTTTCCTCGGTATGGAGTC
genes	hprt	100.95	137	CATGGACTCATCTTGGACAGGACAGA	GCCTTGATGTAGTCCAGCAGGTC
	β-def	102.30	138	GGGAACGAAGATCCAGAGATGCAGTATTGGAC	CTAAGACCTCACAGCACAGCACCTG
Amp genes	hep	104.30	140	CCGTCATGCTCGCCTTCG	CTCACGCATGTAATACGGAATCTTGCATG
	pisc	101.95	111	CAATGATCCATGGGCTTATCC	TTCAGTCTCGCCATTGAAGC

## **Figure**

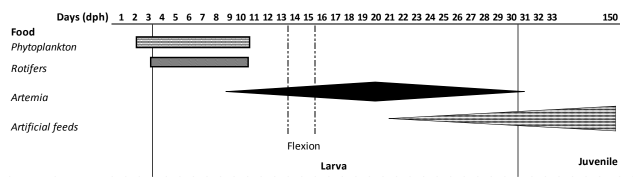
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#### FIGURE CAPTIONS

- 2 Fig. 1. Growth performance of meagre (Argyrosomus regius) larvae. a) The type and relative amount of
- 3 different feed items offered during rearing are shown as an increase or decrease in thickness of the bar
- 4 representing rotifers, Artemia metanauplii and artificial feeds, respectively. b) Total standard length and
- 5 wet weight (mean  $\pm$  SD) during larva rearing, in relation to time (days post-hatching). The regression lines
- 6 describe the best-fit for growth and wet weight curves of meagre larvae.
- 7 Fig. 2. Phylogenetic relationship among defb, hep2, pisc isolated from A. regius obtained by the Maximum
- 8 Likelihood method. The tree with the highest log likelihood (-1611.2745) is shown. The percentages of
- 9 replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are
- 10 indicated at nodes. The tree is drawn to scale, with branch lengths measured in the number of substitutions per
- 11 site.
- 12 Fig. 3. Temporal change in gene expression levels (relative quantification), from larvae, juveniles and
- tissues. from 8-120 dph, of A. regius transcripts coding for defb (a), hep2 (b), pisc (c). Relative expression
- of for *defb*, *hep2*, *pisc* transcripts were normalized using an arithmetic mean of three housekeeping genes:
- GAPDH, HPRT and  $\beta$ -actin. Transcriptional fold changes of *defb*, *hep2* and *pisc* at different time points
- were calculated by comparison to each previous timepoint. Different letters above the bars indicate
- significant differences among different time points (Tukey's test,  $P \le 0.05$ ). Results are expressed as the
- mean  $\pm$  SEM. The biological samples were for larva (n = 10) and tissues, (n = 8) of gene expression. K =
- 19 kidney, Sp = spleen, Gu = gut, Gi = gill.
- Fig. 4. Relative expression of defb (a), hep2 (b), pisc (c) in different tissues. Fish (n = 8) were injected
- 21 intraperitoneally with poly (I:C), LPS and β-glucan and sampled at 24 h post-injection. GAPDH, was
- 22 employed as an internal reference. Asterisks (\*) mark significant differences between stimulated and
- control groups (Kruskal Wallis test,  $P \le 0.05$ ). Data are means  $\pm$  SEM. K = kidney, Sp = spleen, Gu = gut,
- Gi = gill.
- 25 Fig. 5. Expression patterns of defb (a), hep2 (b), pisc (c) mRNA after PAMP stimulation of cell
- suspensions isolated from different meagre (Argyrosomus regius) tissues, extracted from healthy fish (n =
- 27 6). The cells were stimulated with poly (I:C), LPS and β-glucan, and sampled at different time points post-
- stimulation (4, 12 and 24 h), for RNA extraction and qPCR. GAPDH, was employed as an internal
- 29 reference. Asterisks (\*) mark significant differences between stimulated and control groups (Kruskal
- Wallis test,  $P \le 0.05$ ). Data are means  $\pm$  SEM. K = kidney, Sp = spleen, Gu = gut, Gi = gill.

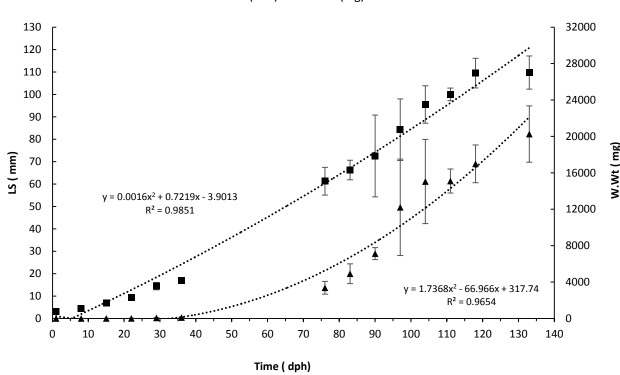
**Fig. 1**.

**a)** 

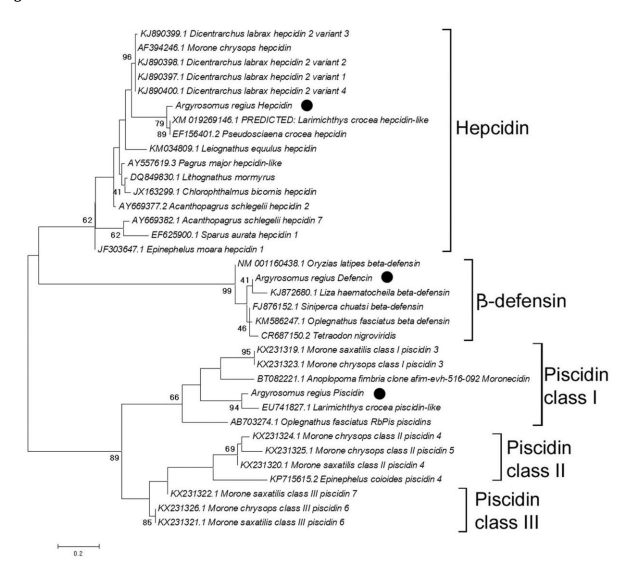


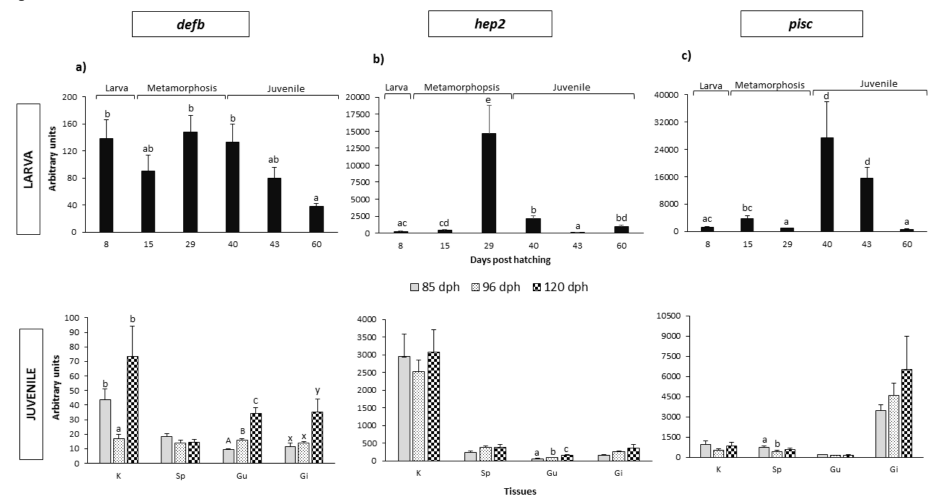
b)





# 42 Fig. 2.





**Fig. 4.** 

