Characterization of Melanin-concentrating Hormone from Olive Flounder (Paralichthys olivaceus)

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The melanin-concentrating hormone (MCH), a cyclic hypothalamic peptide composed of 17 amino acids, was initially identified in chum salmon (Oncorhynchus keta) as a regulator of pigmentation. Mammalian MCHs are cyclic hypothalamic peptides composed of 19 amino acids that regulate food intake and energy homeostasis. The present study examined not only MCH expression of different tissues but also the melanohore aggregation and intracellular Ca2+ influx of fMCH and the other MCH. Real-time qPCR showed that MCH expressed specially in the brain, gonad, and ovary, and expression of MCH was observed during the developmental stages. In the application of synthetic fMCH and both types of synthetic fMCH, dN-fMCH and dC-fMCH, scale melanophore induced significant changes in aggregation activity with various concentrations of MCH. Also, compared to hMCH and sMCH, fMCH exhibited a 36-99.85% increase in relative potency (%), whereas aggregation of dN-fMCH and dC-fMCH remained in a high concentration. However, dispersion was induced rapidly according to low concentration of dN-fMCH and dC-fMCH. We show that fMCH and its derivates were bound human MCHR1 and rat MCHR expressed in HEK293T cells with nano-molar affinity and are likely to be ligand-induced to mobilize intracellular Ca2+. These results may provide new ligands for binding assay with MCHew ligands, as a structure similar to the mammalian MCH structure was discovered in fish. Once the fMCH receptor system is in place, it can be compared to the MCH system of mammals in terms of MCH function.

Key words: Fluorescent imaging plate reader, melanin-concentrating hormone, melanin-concentrating hormone receptor, Paralichthys olivaceus, quantitative real-time PCR

Introduction

Melanin-concentrating hormone (MCH) is a cyclic peptide succeeded in isolating and first characterized in teleosts as a neuropeptide hormone, which involved in aggregation of melanohore in the skin [35]. MCH isolated from the pituitary of chum salmon, Oncorhynchus keta is composed of 17 amino acids (aa), and later the primary structure of MCH has been discovered in coho salmon, O. kisutch [27], chinook salmon, O. tshawytscha [24], tilapia, Oreochromis mossambicus [14], rainbow trout, O. mykiss [6], bonito, Katsuwonus pelamis [19], and recently barfin flounder, Verasper moseri [36] in teleosts. These MCHs showed identical sequences and structure to amino acids, and the only difference which is observed from barfin flounder MCH (Fig. 1) is one variation which has Asn instead of Thr at the N-terminal region. Compared with fish, the sequences and structure of mammalian MCH with consisting of 19 aa also has been confirmed from mice [9], rats [26], and human [28]. Mammalian MCH peptides has an identical cyclic structure and sequence, and as these have two additional and four substitutive amino acids relative to chum salmon MCH, subsequent study of mammalian MCH show that these peptides has been well conserved from vertebrate evolution [25].

MCH peptide with expression and posttranslational cleavage of preproMCH gene encoding MCH has a principal function that regulates the skin color for background adaptation and modulates the release of Adrenocorticothorpe hormone (ACTH), melanin-stimulating hormone (MSH), and somatolactin [5, 7, 14]. Recently, fish MCH plays a role in the regulation of feeding behavior, unlike in rodents, showing that its anorexigenic function is induced by inhibited food intake [30]. These results indicate that the anorexigenic action of MCH with having orexigenic function in the mammals is
mediated by the alpha-MSH signaling pathway. The facts which come to reveal from the mammals MCH is which mammals MCH has the roles as regulation of food intake [1, 11, 13, 22, 29, 30, 33] and exhibits additional regulation functions as behavior of stress, reproduction, grooming, locomotion, and aggression [4, 12, 23, 32].

MCH as orexigenic neuropeptide in mammals mediated signals via a G-protein coupled receptor (MCHR), and two types of MCHR (MCHR1 and MCHR2) was identified until currently. MCHR1 in subtypes was confirmed by the reverse pharmacology study with facts that the natural ligand of the orphan SLC-1 receptor was MCH in mammals [3, 10, 31, 34]. MCHR2 was confirmed in a human genomic sequence with MCHR1 data [17]. Two subtypes of Fish MCHR also have been identified from zebrafish and fugu in whole genome shotgun database, and zebrafish has MCHR1 to be composed of two kind types (MCHR1a and MCHR1b) [21]. Recently, MCHR1 and MCHR2 from the barfin flounder, *Verasper moseri* was identified, and the results that MCHR2 mediates the effect of MCH to control body color for background adaption in the barfin flounder was known in addition to expression feature of MCHR1 [35].

In view of the functions and characterizations relationship between MCH and MCH receptor, It would be of interest to investigate the effects of MCH on melanophore aggregation, signal pathway via MCHR mediation, and the discovery of new ligand for MCH receptor. Consequently, from present work, in order to confirm fMCH characterization using information of the MCH clone first identified from brain cDNA library of Olive flounder, Paralichthys olivaceus in a previous study [16], We compared first amino acid diversity between Olive flounder MCH (fMCH) peptide and different fishes and mammals MCH. Second, Real time qPCR for specific expression of MCH was carried out in various tissues of Olive flounder and embryonic samples of developmental stages. Third, we investigate that melanophore aggregation of fMCH in a Olive flounder scales using synthetic fMCH, hMCH, sMCH, and fMCH derivatives (dN-fMCH; N-terminal deletion, dC-fMCH; C-terminal deletion), and finally, in order to prove using capability as a new ligand of fMCH and to measure the degree of fMCH action, we examined intracellular Ca²⁺ mobilization test by fMCH and different MCHs in human MCHR1/HEK-293T cell and rat MCHR/HEK293T cell with FLIPR (Fluorometric Imaging Plate reader) assay system.

**Materials and Methods**

**MCH Peptide**

25 aa fMCH derived from MCH precursor protein of Olive flounder, salmon MCH (sMCH), human MCH (hMCH), and fMCH derivates as dN-fMCH of 23 aa that the two amino acids were eliminated in amino-terminus and dC-fMCH of 21 aa that the four amino acids were eliminated in carboxy-terminus were synthesized and its quality was checked by a high-performance liquid chromatography (SHIMADZU, Japan). These synthetic fMCH peptide and derivates used in experiments were purchased from the Peptron (Daejon, korea). Synthetic MCHs were dissolved in 50% DMSO, and stored at -70°C until used in FLIPR experiment and pigmentation assay (Table 1).

**Quantitative real-time PCR analysis**

Total RNA of olive flounder with about 150 g was extracted from each 15 tissues (gonad, tail fin, fin, intestine, spleen, heart, skin, stomach, eye, brain, gill, ovary, muscle, kidney and liver) of 100 mg and tissues extraction were ground using mortar and pestle by mixing in liquid nitrogen, and tissues powders disrupted by adding Buffer RLT Plus. After that, the lysates were placed to gDNA Eliminator spin column for removal of extra genomic DNA, and flow-through harvested through the procedure were mixed with 70% ethanol and bound to RNasea spin column. The rest procedure of washing and total RNA elution were carried out according to the manufacturer’s instructions of RNeasy Plus Mini kit (Qiagen, Valencia, CA).

<table>
<thead>
<tr>
<th>Peptide name</th>
<th>Sequence</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>fMCH</td>
<td>H-DTDFN MLRC IGRVY RPCWE SSnip-OH</td>
<td>[25aa] AF236090</td>
</tr>
<tr>
<td>sMCH</td>
<td>H-DTMRM MVGRV YRPCW EV-OH</td>
<td>[17aa] 0910249A</td>
</tr>
<tr>
<td>hMCH</td>
<td>H-DDFM RLMLG RVYRP CWQV-OH</td>
<td>[19aa] AAA63214</td>
</tr>
<tr>
<td>dN-fMCH</td>
<td>H-DNML RCMI MVYRP CWESS NIP-OH</td>
<td>[23aa] AF236090-modified</td>
</tr>
<tr>
<td>dC-fMCH</td>
<td>H-DTNFNL MLRMI IGRVY RPCWE S-OH</td>
<td>[21aa] AF236090-modified</td>
</tr>
</tbody>
</table>

(All peptides were synthesized from Peptron, Korea)
Random hexamer (Bioneer, Korea)-primed cDNA was synthesized from each total RNA, and synthesis was carried out using superscript™ II RNase H− reverse transcriptase (Invitrogen, USA) in a reaction mixture containing 100 pmol of random hexamer (Bioneer) and using 100 ng of total RNA (as measured by the nanodrop, USA) in a final volume of 20 μl, and all reactions were run at 42°C for 60 min and then inactivated at 75°C for 10 min according to the manufacturer’s instructions.

The mRNA levels of MCH in the 15 tissues of Olive flounder were evaluated by Quantitative real-time PCR. These Amplification of qPCR were performed using equal amounts of 80 ng random hexamer (Bioneer, Korea)-primed cDNA as template, a set of forward/reverse primers (1 μM each), QuantiTect™ SYBR® Green PCR kit(Qiagen). The PCR reactions were performed in an DNA Engine Opticon 2 Real-Time PCR Detection System (Bio-rad) at 95°C for 15 min and then 45 cycles of 95°C for 15 sec, 64°C for 20 sec, 72°C for 25 sec based on mchRQ-F, mchRQ-R primers of the Table 1, and the thermocycling protocol for melting curve was extended by heating PCR reaction samples from 65°C to 95°C in 0.2°C increments with a dwell time at each temperature of 1 sec. Real time qPCR of total cDNA in each of the samples was made using a PCR approach specific for cDNA coding for fMCH precursor protein which produced a 331 bp PCR product with primer combination, mchRQ-F primer (forward) 5’-ATG GCC TGA GTT CCT TAC TGG GTG AGC-3’ and mchRQ-R primer (reverse) 5’-CCA GCA GGG TCG GTA CAC TCG TCC TAT-3’ at the PCR condition mentioned above [28].

Melanophore aggregation test with fMCH and MCHs

Olive flounder (Paralichthys olivaceus) was bought from a local fisheries products market (Busan, Korea). For bioassay of MCH, skins were removed from the dorsal surface of Olive flounder of about 150 g, and were harvested by scraping firmly with knife. The skins were then fully soaked in Tyrode Ringer solution containing 0.1 mM phenotamine saline and kept the room temperature throughout the assay. After 30 min pre-incubation period at 20°C, the melanin granules of melanophores on the skins become fully dispersed, and then, serial diluted MCH (1 μM - 2 pM) was added into it and incubated in 30 min for aggregation test of melanophores. Melanophores were photographed by a digital camera (PowerShot A620, Canon, Japan) under phase-contrast microscopy (ITM-400, Nikon, Japan), and the size of concentrated melanophores in photographs was estimated using a freeware program, Scion Image (Scion Corporation, ML, USA), and aggregation degree was measured by relative area of melanophores in each MCH.

Stable cell line

The Stable cell lines which were used in the following FLIPR experiments were obtained from Dr. Olivier Civelli (Department of Pharmacology, University of California, Irvine, California 92612). HEK293T cell lines which human MCHR1 gene are been expressed stably, and HEK293T cell line which rat MCHR gene is been expressed stably were experimented in the dose esponse reactions of Ca2+ intracellular influx.

Measurement of intracellular Ca2+ concentration

hMCHR1/HEK293T cell, hMCHR2/HEK293T cell and ratMCHR/HEK293T cell were cultivated until 80-90% confluence and about 90,000 cells per well were inoculated in lysine-coated 96 well FLIPR plate and Those were cultivated for 18 hr at 37°C. Supernatant was discarded and The cells were loaded with Fluo-4 dye (Molecular probe) in standard buffer solution (130 mM NaCl, 2 mM CaCl2, 5 mM KCl, 10 mM glucose, 0.45 mM KH2PO4, 0.4 mM Na2HPO4, 8 mM MgSO4, 4.2 mM NaHCO3, 20 mM HEPES and 10 mM probenecid) with 0.1% fetal bovine serum for 1 hr at 37°C, then washed with a standard buffer solution. In 96 well plate, hMCH, sMCH, fMCH and fMCH derivative was added at different concentration and Drug Plate were completed. And measurement of Ca2+ influx changes was obtained from FLIPR system (Fluorometric Imaging Plate Reader; Molecular Devices) in 96-well plates at 488 nm for 210 sec.

Statistical analysis

The EC50 values for MCH were obtained from sigmoidal fits using a nonlinear curve-fitting program (Prism version 4.0, GraphPad Software, La Jolla, CA, USA). Melanophore aggregation assay were performed in ten replicates and Ca2+ influx changes with MCHs in the HEK293T cell lines were performed in triplicate. These data were expressed throughout sigmoidal dose-response using 95% confidence intervals. Specially, the equation of melanophore aggregation is Sigmoidal dose-response as Y is Bottom + (Top-Bottom)/ (1+10(α((LogEC50-X)))). Also, fMCH expression data was expressed with mean ± standard deviation (n=3).
Results

Diversity of fMCH and its derivatives, sMCH, hMCH and barfin flounder MCH

Olive flounder MCH (fMCH) which was confirmed initially from previous paper [16] was revealed to contain mature form of hormone with 25 amino acid residues. The fish MCH and mammals MCH compared with fMCH, which becomes known already was had the identical cyclic structure, but were showing wide difference from amino acid sequence (Fig. 1). A comparative structural analysis was revealed that fMCH had a relatively high identity of 78% to mammals MCH, compared with the fish which was kept 70% identity, and identity between fMCH and barfin flounder MCH also put out 70%, and was similar to other fishes. And then, fMCH of the cyclic structure which almost agrees to different MCH will be functional characteristic in the melanophore aggregation and Ca\(^{2+}\) mobilization, and may be important to novel MCH in the following experiment.

Tissue and developmental expression of MCH in Olive flounder

In order to confirm the presence of MCH, Real-time qPCR was performed with cDNA samples prepared from the total RNA of the respective tissues and various developmental stages after fertilization (Fig. 2). The expression level of fMCH with replaced at value of \(\log_{10}[\text{fMCH expression (ng)}]\) in various tissues of Olive flounder showed that it was expressed specifically only in the brain, gonad, and ovary but not in other tissues to be test (Fig. 2A), especially, expression level of fMCH was significantly 2 more times (2.74±0.15) greater in the brain tissue than in gonad (1.00±0.46) and ovary tissues (1.03±0.04), and no difference almost was observed in the expression levels of fMCH between the ova-

![Fig. 1. Comparison of amino acid sequences of fish and mammals MCHs. The two cysteine residues are line to form a cyclic structure with disulfide bonds. Residues which differ from fish MCH sequence are underlined. sMCH, salmon/tilapia MCH; hMCH, barfin flounder MCH; hMCH, rat/mouse/human MCH; fMCH, Olive flounder MCH.](image)

![Fig. 2. Quantitative real-time PCR analysis of fMCH transcripts in different tissues and in developmental stages. 14 tissues (A) and samples (B) of multiple time zone after fertilization was used in a quantitative real-time PCR and amplified with primers specific for fMCH gene. fMCH expression quantity was calculated with replaced at value of \(\log_{10}[\text{fMCH expression (ng)}]\). The level of fMCH with fMCH expression amount was represented with mean ± standard deviation (n=3).](image)
ry and gonad. fMCH expression in the developmental stages of the fertilized flounder eggs were observed from 30 hr after fertilization to 80 hr after fertilization, even though IMCH expression was not observed from fertilization time to 30 hr after fertilization time, and the expression quantity of fMCH which when from 37 hr (1.46±0.02) went in 80 hr (2.79±0.59) increased progressively.

### Melanophore aggregation test

Melanophore in vitro dose responses to various concentrations (1 μM - 2 pM) of MCHs were measured from the Olive flounder (Paralichthys olivaceus) scale in order to confirm relative aggregation activity of fMCH (Fig. 3). In the scale melanophores showed full dispersion after immediately removal from the Olive flounder, when high concentration of 1 μM MCHs correspond to 10^-6 M was added in scale, those were fully aggregated increased with the MCHs except sMCH (Fig. 3A), and also photograph showing effect of MCHs was relatively weak aggregation activity in a sMCH (Fig. 3B). Aggregation activity from MCHs where increased progressively put out a serious difference. When fMCH of 10^-6-10^-9 M was applied to the scale preparation, degree of aggregation was continued and maintained from concentration of wide phase. fMCH derivate with N-terminal (dN-fMCH) and C-terminal deletions (dC-fMCH), and hMCH revealed which melanophores resulted in almost dispersion in compliance with the concentration which increased with 10^-7M, although dN-fMCH, dC-fMCH and hMCH led to rapid aggregation at a high concentration of 10^-6 M. EC50 values exhibiting concentration of MCHs causing half maximal aggregation of scale melanophores was lowest from fMCH of 0.16nM concentration, moreover as this result, the relative potency ranking of the MCHs was as follows: fMCH > sMCH > dN-fMCH > hMCH > dC-MCH, fMCH proved most responsive to the melanophore of Olive flounder (Table 2).

These results suggest the fact that fMCH aggregation activity was highest from the melanophore of Olive flounder as a result of EC50 value and relative potency of MCHs (Table 2), and fMCH will come to new counterpart that regulates skin coloration in fish.

### Ca²⁺ mobilization

To confirm if the ligand function of neuropeptide by intracellular Ca²⁺ mobilization as well as the aggregation effect of fMCH is mediated by MCH receptor, The binding characteristics by intracellular Ca²⁺ influx of fMCH and the other MCHs were carried out with HEK293T cells stably expressing human MCH-R1 and rat MCH-R. fMCH and the other MCHs induced Ca²⁺ influx in the hMCH-R1/HEK293T cells, and the MCHs concentration to induce a half-maximum response (EC50) were 4.7~10.0 nM, indicating which MCHs are an affinity ligand for human MCH-R1 (Fig. 4A). Especially, fMCH, which has the lowest degree of EC50, activated effec-

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Table 2. Concentrations of MCH peptides causing half maximal aggregation of scale melanophores of the Olive flounder, and in vitro relative potencies as compared to fMCH

<table>
<thead>
<tr>
<th>Substance (aa)</th>
<th>EC50 (nM)</th>
<th>Relative potency</th>
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</thead>
<tbody>
<tr>
<td>fMCH [25aa]</td>
<td>0.16</td>
<td>1.00</td>
</tr>
<tr>
<td>sMCH [17aa]</td>
<td>0.25</td>
<td>0.64</td>
</tr>
<tr>
<td>hMCH [19aa]</td>
<td>54</td>
<td>0.0029</td>
</tr>
<tr>
<td>dN-fMCH [23aa]</td>
<td>19</td>
<td>0.0084</td>
</tr>
<tr>
<td>dC-fMCH [21aa]</td>
<td>103</td>
<td>0.0015</td>
</tr>
</tbody>
</table>
tively human MCHR1 with an EC₅₀ of 4.7 nM compared to hMCH. Moreover, because the fMCH was having the structure which is nearer in the hMCH than sMCH, fMCH together with hMCH will become as the different new ligand. Experiments of MCHs dose responded to rat MCHR showed results similar to hMCHR1/HEK293T cells test, that fMCH, dN-fMCH, and dC-fMCH are ligand having relatively high potency than hMCH and sMCH as induced a high dose response (Fig. 4 B). dN-fMCH and dC-fMCH were confirmed by more affinitive MCH than hMCH for rat MCHR with EC₅₀ of 2.0~1.5 nM.

These result suggest that fMCH is likely to be an endogenous ligand for human MCHR1 and rat MCHR because this peptide led to a relative high potency of 48% than hMCH responded to hMCHR1 and to a relative high potency of 33% than hMCH responded to rat MCHR, and dN-fMCH and dC-fMCH had a possibility with the candidate of a potent agonist for human MCH-R1 and rat MCHR.

Discussion

Amino acid sequence of fMCH described in previous study was longer in Olive flounder than in a different fishes and mammals. fMCH had eight additional and five substitutive amino acids (Fig. 1) compared with fishes and had six additional and four substitutive amino acids compared with mammals, but sequence inside a cyclic structure formed by a disulfide bond was identical to fishes and mammals except on replacement. One of them in the cyclic structure, Ile¹¹ in Olive flounder and Val7 in fishes and Leu¹ in mammals, was a physicochemically conservative replacement. Excepts the fact that sequence size is long region which are essential for binding to the receptor is identically compose of as Arg⁴ residue in the N-terminal sequence and Trp¹³ residue in the C-terminal sequence, then this is more conserved in the different MCH region. The other four additional amino acids located in outside N-terminal sequence and C-terminal sequence compared to fishes MCH do not accurately revealed those functions, but it is likely that the rest region of fMCH may have the function which is not become known until now.

In Olive flounder, fMCH expression in level of mRNA is very unique, and fMCH is only expressed specifically in brain, gonad and ovary. This showed that fMCH expression appeared similar to salmon MCH [27, 28] and tilapia MCH study [14], then these reports are the same as there is expressed at the pituitary and hypothalamus of brain. IMCH expression in gonad and ovary, which showed weak expression than brain tissue, has very doubtful point, but in compliance with report [15], MCH neurones are detectable only in reproductively mature animal as the telencephalon, suggesting a potential role in reproduction and/or behavior, then fMCH expression also will be possible specifically in gonad and ovary as well as brain like mammals MCH.

The identity of structure was made to demand an experiment, and aggregation activity of fMCH in scale melanophor of Olive flounder is advanced and getting the answer which is clear. The administration of fMCH, hMCH, sMCH, dN-MCH and dC-MCH induced melanophore aggregation in a dose dependent manner (Fig. 3A, Fig. 3B). This result, which the fMCH peptide showed the highest potency of aggregation activity in the Olive flounder scale and revealed the lowest EC₅₀ value (Table 2, 0.16 nM), are comparable to others. EC₅₀ as the potency of aggregation activity of MCH
varies in various teleost fish with value between 30 pM and 1 nM [23, 35, 36]. This result suggests that fMCH will be worth of neuropeptide first identified in the Olive flounder, even though the range of EC50 value might be explained by the trivial and technical difference in the bioassays used. Derivatives of fMCH, which lack the N- and C- terminal region changed in native fMCH have reported to have diminished aggregation activity (when tested on melanophore, although they were able to induce virtually complete aggregation in a high concentration (10^8 M, Fig. 3B). But derivatives activity in Ca^{2+} influx test with FLIPR assay system show obvious similarity in their native fMCH and is recognized as new ligand enough in activation of MCH-receptor.

MCH and its receptors play an important part in the complex network involving peripheral and central signals regulating food intake and energy expenditure [18]. In intracellular Ca^{2+} influx of fMCH, dN-fMCH and dC-fMCH, although fMCH EC50 shows the value which is relative to sMCH and hMCH, there shows higher value. Furthermore, these activate the rat MCHR that there is not rat MCHR2 as well as ligand activating human MCHR1, fMCH could be accomplish a role human and rat MCH receptor and the other function. This may suggest that fMCH and derivatives functions not only melanophore aggregation but also food intake showing in mammals.

In conclusion, we suggest that first fMCH structure was more similar to mammals than fishes during the diversity analysis between and fishes MCH and mammals MCH, furthermore fMCH amino acid sequence conserved with MCHs of existing. Second, specific expression of MCH mRNA was observed in the different tissues, gonad and ovary, excepts brain. This result show that there is indicating variety of fMCH expression as rodent MCH expression profile. Third, our data exhibit the new type of MCH possible involved with melanophore aggregation activity, and peptide that there is a value which will research in fishes identified from Olive flounder, Paralichthys olivaceus. Finally, we conclude that nanomolar concentrations of fMCH activate human MCHR1 and rat MCHR related to signaling pathway system, and that could be as higher degree of ligand compared to other MCHs.

Acknowledgement

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References


초록: 양식넙치 멜라닌 농축 호르몬의 특성

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멜라닌 농축 호르몬(melanin-concentrating hormone, MCH)은 17개의 아미노산으로 구성된 환형의 시상하부 쵸터드로 색소 침착의 조절인자로서 연어에서 처음 분리되었다. 포유동물의 MCH는 19개의 아미노산으로 구성되어 있으며 식사 및 에너지 향상을 조절하는데 관여한다. 본 연구에서는 양식넙치의 다양한 조직에서 MCH 유전자의 발현 분포, 멜라닌 촉매 세포의 집적, 포유동물 MCH 수용체와 양식넙치 MCH의 상호작용을 조사하였다. Real-time qPCR을 이용하여 뇌, 정소, 난소에서 MCH 유전자의 발현이 나타나는 것을 확인하였고, 수정 후 발달 단계에서도 MCH 유전자의 발현을 확인할 수 있었다. 합성된 연어 sMCH, 포유류 hMCH, 양식넙치 fMCH, dN-fMCH, dC-fMCH를 양식 넙치의 표피에 처리했을 때 다양한 농도에 따라 멜라닌 함유 세포의 집적이 다양하게 나타났다. 연어 sMCH, 포유류 hMCH에 비해 양식넙치 fMCH의 멜라닌 함유세포의 집적도가 36~99.85%로 비슷한 값을 나타냈으나 양식넙치 dN-fMCH, dC-fMCH를 처리한 경우 양식넙치 fMCH에 비해 높을 농도에서 집적이 나타나고 짧은 시간에 분산되었다. 또한, 인간 MCH 수용체가 양식넙치 MCH 수용체가 발현된 포유동물의 세포주와 양식넙치 fMCH를 처리하여 각 수용체와 결합하는 것을 확인하였다. 이러한 결과는 어류에서 발현되는 MCH가 포유동물의 MCH와 유사한 구조를 가지고 있어 MCH 수용체에 대한 새로운 리간드로서 제공될 수 있으며, 향후 어류의 MCH 수용체에 적용할 수 있을 것이다.