



Journal of
**Fisheries and
Aquatic Science**

ISSN 1816-4927



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Corrigendum

“Expression Levels of Hormone Receptor and Vitellogenin mRNAs in Livers of Thai Medaka, *Oryzias latipes*, inhabiting the suburbs of Bangkok, Thailand” published in Journal of Fisheries and Aquatic Science 6 (4): 438-446, 2011

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Authors regretted that the errors by author were found into this article. Now on the request of author Page No. 439 and 440 has been changed due to some errors by author that Page No. 439 and 440 republished as under:

On page 439

In line 25, for MATERIALS AND METHODS, correction is “Fish: Adult Thai medaka were captured in ponds and ditches in the Nakhonpathom and Ratchaburi provinces which are suburbs of Bangkok, Thailand, during February through to April of 2010. The standard length of the captured Thai medaka was 14-16 mm. Average water temperature in those areas was $28 \pm 1^\circ\text{C}$ ”.

On page 440

In line 2, correction is “Thai medaka were collected from February through to April because of the beginning of dry season, but it was hard to consider as the breeding or non-breeding season for this species. Then, conducting experiments might be included both conditions”.

In line 9, correction is “Total RNA from each fish sample was extracted using the RNeasy Mini Kit (Qiagen) according to the manufacturer’s protocol and then treated with DNase1 (Takara, Tokyo, Japan) for 30 min at 37°C ”.

In line 15, for semi-quantitative RT-PCR, correction is “The primers used to amplify androgen receptor (AR) were 5'-GTCGTGGATGGGGGTGATGGTG-3' and 5'-CCGACTGGAGGTAGTCCAGCAGC-3' designed from the nucleotide sequences of teleosts in GenBank sequence database”.

In line 23, correction is “Amplification of cDNA was conducted using the following cycling conditions: 95°C for 30 sec for the denaturation step; 55°C (AR and Vtg), 56°C (ER) or 58°C (β -actin) for 1 min for the annealing step and 72°C for 1 min for the extension step”.