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#### INVESTIGATING THE GENETICS OF SEX DETERMINATION IN EUROPEAN SEA BASS (Dicentrarchus labrax) USING RAD-SEQ

# Palaiokostas C.<sup>1</sup>, Bekaert M.B.<sup>1</sup>, Taggart J.B.<sup>1</sup>, Gharbi K.<sup>2</sup>, McAndrew B.J.<sup>1</sup>, Chatain B.<sup>3</sup>, Penman D.J.<sup>1#</sup>, Vandeputte M.<sup>3,4</sup>

 <sup>7</sup> Institute of Aquaculture, School of Natural Sciences, University of Stirling, Stirling FK9 4LA, Scotland, United Kingdom
<sup>2</sup> Edinburgh Genomics, Ashworth Laboratories, King's Buildings, University of Edinburgh, Edinburgh EH9 3JT, Scotland, United Kingdom
<sup>3</sup> Ifremer, Chemin de Maguelone, 34250 Palavas-les-Flots, France
<sup>4</sup> INRA, UMR1313 Génétique animale et Biologie intégrative, 78350 Jouy-en-Josas, France

European sea bass (*Dicentrarchus labrax*) is one of the most important farmed species for Mediterranean aguaculture. The observed sexual growth and maturity dimorphism in favour of females adds additional value to deciphering its sex-determining system. Current knowledge indicates the existence of a polygenic sex-determining system interacting with temperature. Restriction-site Associated DNA (RAD) sequencing was used in a test panel of 175 offspring originating from a F2 factorial cross between two dams and four sires from one F1 family between parents with contrasted sex tendencies. In total, 1.156,659,542 raw reads (100 bases long) were produced (578,329,771 paired-end reads). After removing low quality sequences (quality score under 30), ambiguous barcodes and orphaned paired-end reads, 76.7% of the raw reads were retained (886,927,866 reads) from which 56,696 unique RADtags were retrieved. In order to maximise the number of informative markers and minimise the amount of missing or erroneous data, we used RAD-tags retrieved in at least 75% of the samples, and carrying one or two single nucleotide polymorphisms (SNPs). The family structure was identified *a posteriori* using R/hsphase and Vitassign with the SNP markers from the offspring and parents. A consensus SNP-based linkage map was constructed, consisting of 5,097 SNPs grouped into 24 linkage groups. Indications for putative sex-determining OTL are provided in linkage groups 13, 19 and 21. A preliminary study was conducted testing whether breeding values, estimated by including the additive SNP effects, could be used as a predictive measure of phenotypic sex. Prediction of phenotypic sex within each of the eight full-sib families, using genomic models (MCMC-BLUP, BayesCPi), consistently outperformed pedigree BLUP. The advantage of using genomic models was particularly evident in dam half-sib family 2, where one of the sex-determining QTLs had been detected. Altogether, our results further support the polygenic hypothesis for sex determination in sea bass, while they also highlight the potential use of within-family genomic selection to rank individuals within families for their estimated breeding value for sex tendency.

Keywords: RAD-seq, Dicentrarchus labrax, QTL, Genomic Selection

<sup>§</sup> Corresponding author. Tel.: + 44 (0)1786 467901; Fax: + 44 (0)1786 472133 E-mail address: d.j.penman@stir.ac.uk