

1 **Corticosteroid and progesterone transactivation of** 2 **mineralocorticoid receptors from Amur sturgeon and tropical gar**

3
4 Akira Sugimoto¹, Kaori Oka¹, Rui Sato¹, Shinji Adachi²,
5 Michael E. Baker^{3,*}, Yoshinao Katsu^{1,4,*}

6
7 ¹Graduate School of Life Science, Hokkaido University, Sapporo, Japan; ²Faculty of Fisheries
8 Sciences, Hokkaido University, Hakodate, Japan; ³Department of Medicine, University of
9 California, San Diego, CA, USA; ⁴Department of Biological Sciences, Hokkaido University,
10 Sapporo, Japan

11
12 *Corresponding authors:

13 Michael E. Baker: mbaker@ucsd.edu

14 Yoshinao Katsu; ykatsu@sci.hokudai.ac.jp

15 16 **Abstract**

17 The response to a panel of steroids by the mineralocorticoid receptor (MR) from Amur sturgeon
18 and tropical gar, two basal ray-finned fish, expressed in HEK293 cells was investigated.

19 Half-maximal responses (EC50s) for transcriptional activation of sturgeon MR by
20 11-deoxycorticosterone, corticosterone, 11-deoxycortisol, cortisol and aldosterone, and
21 progesterone were between 13 pM and 150 pM. For gar MR, EC50s were between 8 pM and 55
22 pM. Such low EC50s support physiological regulation by these steroids of the MR in sturgeon
23 and gar. Companion studies with human MR and zebrafish MR found higher EC50s compared
24 to EC50s for sturgeon and gar MR, with EC50s for zebrafish MR closer to gar and sturgeon MR
25 than was human MR. For zebrafish MR, EC50s were between 75 pM and 740 pM; for human
26 MR, EC50s were between 65 pM and 2 nM. In addition to progesterone, spironolactone and
27 19nor-progesterone were agonists for all three fish MRs, in contrast to their antagonist activity for
28 human MR, which is hypothesized to involve serine-810 in human MR because all three steroids
29 are agonists for a mutant human Ser810Leu-MR. Paradoxically, sturgeon, gar and zebrafish MRs
30 contain a serine corresponding to serine-810 in human MR. Our data suggests alternative
31 mechanism(s) for progesterone, spironolactone and 19nor-progesterone as MR agonists in these
32 three ray-finned fishes and the need for caution in applying data for progesterone signaling in
33 zebrafish to human physiology.

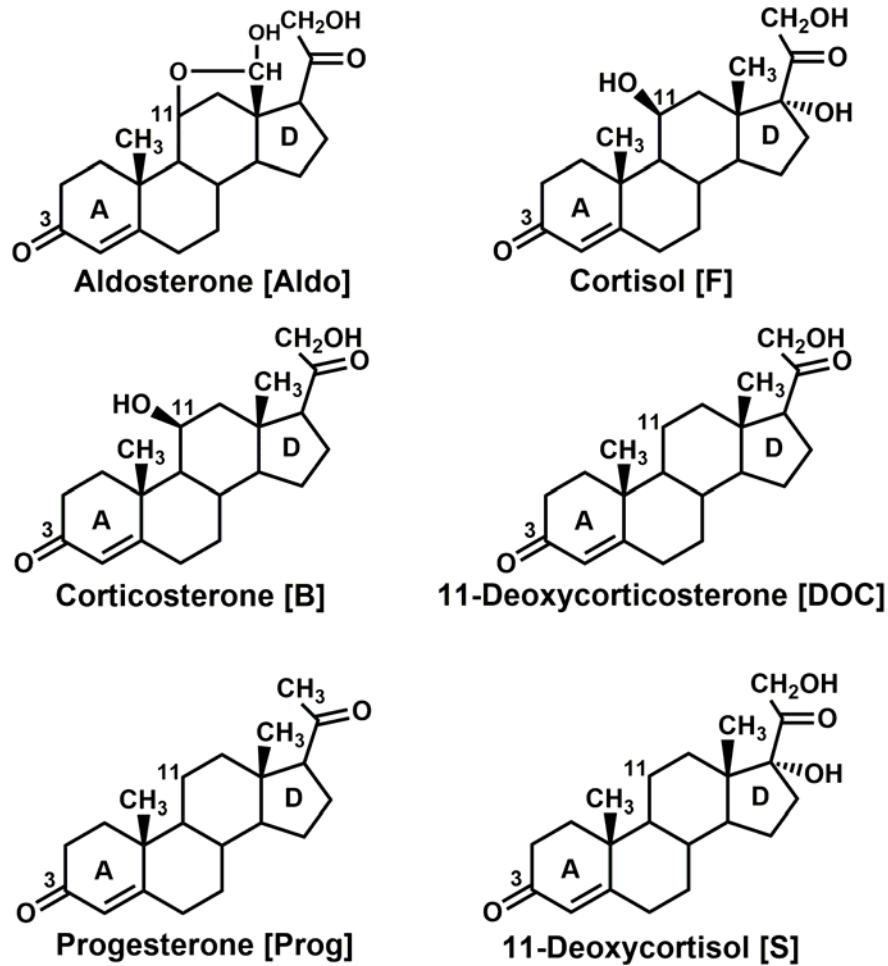
1 INTRODUCTION

2 The mineralocorticoid receptor (MR) is a transcription factor that belongs to the
3 nuclear receptor family, a diverse group of transcription factors that also includes
4 receptors for androgens (AR), estrogens (ER), glucocorticoids (GR) and progestins (PR),
5 and other small lipophilic ligands, such as thyroid hormone and retinoids [1-5]. The
6 MR and GR are descended from a common corticosteroid receptor (CR), which are
7 present in lampreys and hagfish [5-7]. Several corticosteroids (Figure 1), including
8 aldosterone (Aldo), cortisol (F), 11-deoxycortisol (S), corticosterone (B) and
9 11-deoxycorticosterone (DOC), as well as progesterone (Prog), are transcriptional
10 activators of Atlantic sea lamprey CR and hagfish CR [6]. Among these steroids, Aldo,
11 the main physiological activator of the MR in human and other terrestrial vertebrates
12 [8-11], had the lowest half-maximal response (EC50) for transcriptional activation of
13 the CR. This strong response to Aldo is surprising because Aldo is not found in either
14 lamprey or hagfish serum [6]. S, which along with DOC is present in Atlantic sea
15 lamprey serum, has been found to have mineralocorticoid activity in lamprey [12, 13].

16 Distinct MR and GR genes first appear in cartilaginous fishes (Chondrichthyes),
17 such as sharks, rays and skates [6, 14]. Carroll et al. [14] determined EC50s of several
18 corticosteroids for skate MR; EC50s were 70 pM for Aldo, 30 pM for DOC, 90 pM for
19 B, 1 nM for F and 22 nM for S. In teleosts, which comprise about 95% of known
20 ray-fish species (*Actinopterygii*), corticosteroid activation of the MR has been
21 investigated for cichlid [15], trout [16], carp [17], midshipman fish [18] and zebrafish
22 [19], with Aldo, F and DOC being the principal steroids that were studied. Although
23 Aldo has not been found in teleost fish [20], Aldo has a low EC50 for teleost MRs,
24 similar to that found for Aldo activation of lamprey CR and skate MR. DOC also has
25 a low EC50 for teleost MRs, and DOC has been proposed as mineralocorticoid in fish
26 [16, 21-25]. F also has been proposed to be ligand for teleost fish MR [22, 24, 25].
27 The response of the teleost MRs to B and S, which are found in fish [25, 26], has been
28 studied only in trout, in which the EC50s are 10 nM for B and 3.7 nM for S [16].
29 Interestingly, spironolactone (spiron) and Prog are agonists for trout MR [16], in
30 contrast to their antagonist activity for human MR [16, 27]. Spiron also is an agonist
31 for zebrafish MR [19]. Together, these studies indicate that several corticosteroid(s)
32 are potential transcriptional activators of teleost MRs [22, 25, 28, 29].

33 An important gap in our understanding of the evolution of selectivity of
34 ray-finned fish MRs for steroids is the absence of data on the MR in Chondrostei
35 (sturgeons, paddlefishes, reedfishes, bichirs) and Holostei (bowfins, gars), which
36 evolved before a fish-specific genome duplication occurred after the split of the

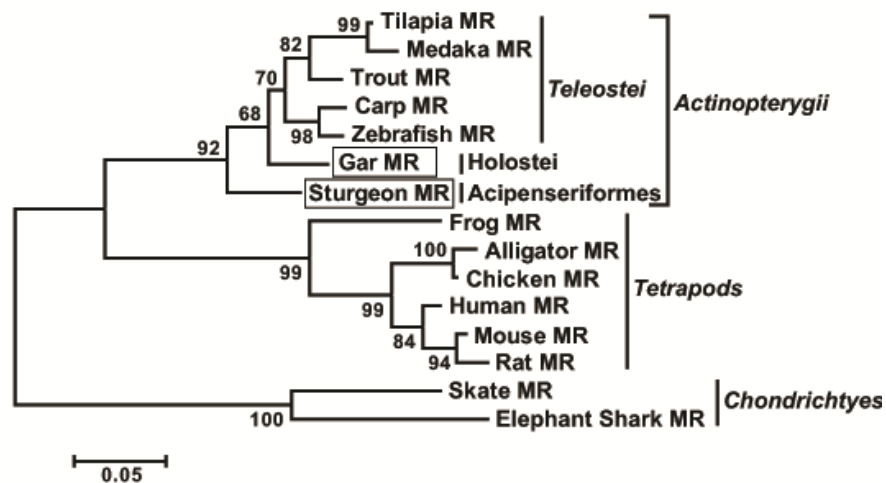
- 1 Acipenseriformes (sturgeons) and the Semionotiformes (gars) from the lineage leading
2 to teleost fish, but before the divergence of Osteoglossomorpha (Figure 2) [30-32].
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Figure 1. Structures of potential steroid regulators of fish MR.

Aldo, the physiological ligand for terrestrial vertebrate MRs, is not found in fish [20]. F and DOC have been proposed to be mineralocorticoids in teleosts [22, 25]. S is a ligand for corticosteroid receptor in lamprey [12]. Progesterone is an antagonist for human MR [27].



1

2 **Figure 2. Phylogenetic relationship of sturgeon and gar MRs to other vertebrates.**

3 To investigate the relationship of sturgeon and gar to other fish, we constructed a phylogenetic
4 tree of the steroid-binding domains on MRs in sturgeon, gar, selected teleosts, elasmobranchs
5 and tetrapods. The phylogenetic tree was constructed using the maximum likelihood with
6 JTT+G model with 1000 bootstrap replications, which are shown as percentages at the nodes of
7 the tree.

8

9 Our interest in the evolution of steroid hormone action [4, 5, 33, 34] prompted us to
10 investigate transcriptional activation of the MR from Amur sturgeon, *Acipenser schrenckii*, and
11 tropical gar, *Atractosteus tropicus* by a broad panel of corticosteroids (Aldo, F, B, DOC, S), Prog,
12 spiron and 19nor-progesterone (19norP), a steroid that has not previously been studied for
13 activation of fish MR. To gain further insight into the evolution of steroid specificity, we
14 compared our results with companion studies of zebrafish and human MRs. In agreement with
15 studies of teleost MRs, we find that Aldo and DOC have the lowest EC₅₀ (highest activity) for
16 sturgeon and gar MRs. However, we also find that S, B, F, and Prog have low EC₅₀s,
17 consistent with these steroids also having a physiological role as ligands for these MRs. Spiron
18 and 19norP also activated sturgeon and gar MRs. In comparison, zebrafish MR has a strong
19 response to Aldo and DOC and a good response to B, F, S Prog, spiron and 19norP, while human
20 MR has strong response to Aldo, DOC and B and a good response to F and S, and a weak
21 response to Prog and 19norP and no response to spiron. The weak response to Prog and 19norP
22 and absence of a response to spiron by human MR is in agreement with other studies [27, 35, 36].
23 The strong response to 19norP, Prog and spiron of sturgeon, gar and zebrafish MR is perplexing
24 because the basis for the low response to to these steroids by human MR is thought to be due to
25 the presence of Ser-810 on α -helix 5 [27, 37, 38]. 19norP, Prog and spiron are agonists for

1 human MR with Ser810Leu mutation [27, 37, 38]. Sturgeon, gar and zebrafish MRs contain a
2 serine corresponding to serine-810 in human MR, suggesting the presence of an alternative
3 mechanism for these steroids acting as MR agonists in these three ray-finned fishes, as well as
4 the need to apply caution in interpreting data on Prog activity in zebrafish to human physiology.

6 **MATERIALS AND METHODS**

7 **Animals and chemical reagents**

8 Amur sturgeon and tropical gar were obtained as described previously [32]. All
9 experimental procedures involving live fish followed the policies and guidelines of the Hokkaido
10 University Animal Care and Use Committee. Aldosterone (Aldo) (CAS 52-39-1),
11 corticosterone (B) (CAS 50-22-6), cortisol (F) (CAS 50-23-7), 11-deoxycortisol (S) (CAS
12 152-58-9), 11-deoxycorticosterone (DOC) (CAS 64-85-7), progesterone (Prog) (CAS 57-83-0),
13 19nor-progesterone (19norP) (CAS 472-54-8), spironolactone (Spiron) (CAS 52-01-7),
14 5 α -dihydrotestosterone (DHT) (CAS 521-18-6), and 17 β -estradiol (E2) (CAS 50-28-2) were
15 purchased from Sigma-Aldrich. For the reporter gene assays, all hormones were dissolved in
16 dimethyl-sulfoxide (DMSO) and the final concentration of DMSO in the culture medium did not
17 exceed 0.1%.

19 **Molecular cloning of mineralocorticoid receptors**

20 Two conserved amino acid regions, GCHYGV and LYFAPD of vertebrate MRs were
21 selected and degenerate oligonucleotides were used as primers for PCR. First-strand cDNA
22 was synthesized from 2 μ g of total RNA isolated from the liver after amplification, and an
23 additional primer set (CKVFFK and LYFAPD) was used for the second PCR. The amplified
24 DNA fragments were subcloned with TA-cloning plasmid pGEM-T Easy vector, sequenced using
25 a BigDye terminator Cycle Sequencing-kit with T7 and SP6 primers, and analyzed on the 3130
26 Genetic Analyzer (Applied Biosystems). The 5'- and 3'-ends of the mineralocorticoid receptor
27 cDNAs were amplified by rapid amplification of the cDNA end (RACE) using a SMART RACE
28 cDNA Amplification kit.

30 **Database and sequence analysis**

31 MRs for phylogenetic analysis were collected with Blast searches of GenBank. A
32 phylogenetic tree for MRs was constructed by the Neighbor-Joining Method [39] after sequences
33 were aligned by MUSCLE [40] using several fish, frog, alligator, chicken, rat, mouse, human
34 MRs. Maximum likelihood (ML) analysis was conducted using the JTT+G model. Statistical
35 confidence for each branch in the tree was evaluated by the bootstrap method [41] with 1000
36 replications. We used the MEGA5 program [42] for these analyses.

1

2 **Construction of plasmid vectors**

3 Full-coding regions of mineralocorticoid receptors were amplified by PCR with KOD
4 DNA polymerase. PCR products were gel-purified and ligated into pcDNA3.1 vector. Mouse
5 mammary tumor virus-long terminal repeat (MMTV-LTR) was amplified from pMSG vector by
6 PCR, and inserted into pGL3-basic vector containing the *Photinus pyralis* luciferase gene. All
7 constructs were verified by sequencing.

8

9 **Transactivation Assay**

10 Human embryonic kidney 293 (HEK293) cells were used in the reporter gene assay.
11 Transfection and reporter assays were carried out as described previously [33], except that we
12 used PEI-max as transfection reagent [43]. All transfections were performed at least three times,
13 employing triplicate sample points in each experiment. The values shown are mean \pm SEM
14 from three separate experiments, and dose-response data and EC50 were analyzed using
15 GraphPad Prism.

16

17 **Statistical methods**

18 Results are presented as mean \pm SE (SEM) from three separate experiments. All
19 multi-group comparisons were performed using one-way ANOVA followed by Bonferroni test.
20 Dose-response data and EC50 were analyzed using GraphPad Prism. $P < 0.05$ was considered
21 statistically significant.

22

23 **RESULTS**

24 **Isolation of mineralocorticoid receptors from sturgeon and gar**

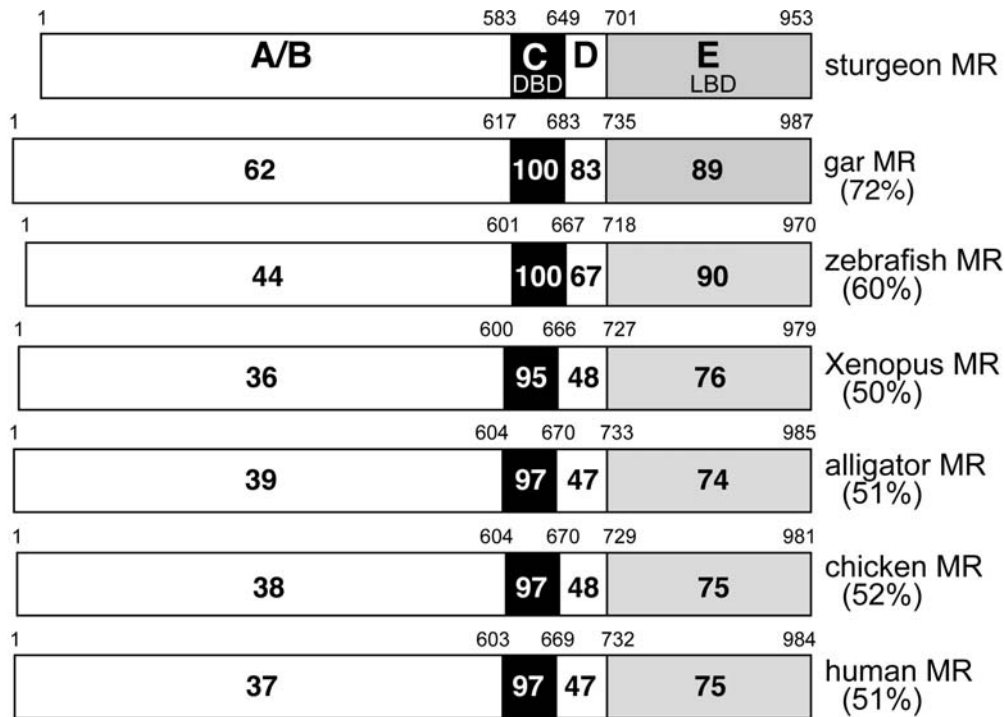
25 We cloned sturgeon MR cDNA containing an open reading frame encoding 953 amino
26 acids (GenBank accession LC149818)], and gar MR cDNA containing an open reading frame
27 encoding 987 amino acids (GenBank accession LC149819). Sturgeon and gar MR sequences
28 can be divided into four domains (Figure 3). The overall amino acid identity between these two
29 MRs was 72%, with particularly high sequence identities for the DBD (100%) and LBD (89%)
30 (Figure 3). Comparison of sturgeon MR with five other species (human, chicken, alligator,
31 *Xenopus*, and zebrafish) revealed that sturgeon MR had identities of 44-36% in A/B
32 domains, 100-95% in DBDs, 67-47% in D domains, and 90-74% in LBDs (Figure 3).

33

34 **Phylogenetic analysis of ancient fish corticoid receptors**

35 To investigate the evolutionary position of gar and sturgeon MR in relationship to other
36 fish MRs and tetrapods, we collected MR sequences from several teleosts, skates and elephant

1 shark and selected terrestrial vertebrates. Consistent with the evolution of Acipenseriformes
 2 and Holostei, phylogenetic analysis places sturgeon and gar MRs close to the base of ray-finned
 3 fish (Figure 2).
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 6 **Figure 3. Comparisons of functional domains in sturgeon, gar, zebrafish, *X. laevis*, alligator,**
 7 **chicken and human MRs.**

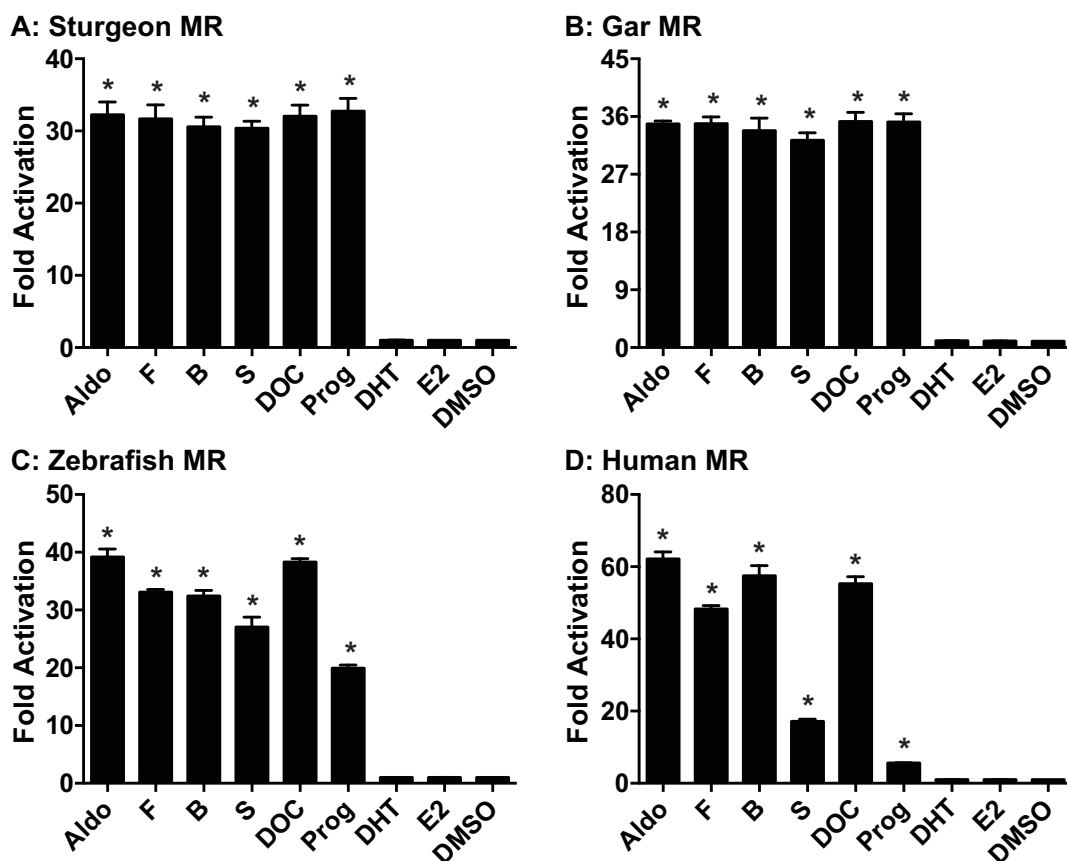
8 Comparison of the domains in sturgeon MR gar, zebrafish, *X. laevis*, alligator, and human MR
 9 MR. The functional A/B domain, C domain, D domain and E domain are schematically
 10 represented with the numbers of amino acid residues at each domain boundary indicated. The
 11 percentage of amino acid identity between domains is depicted. GenBank accession numbers
 12 are: LC149818 for sturgeon MR; LC149819 for gar MR; NM_001100403 for zebrafish MR;
 13 NM_001090605 for *Xenopus* MR; AB701406 for alligator MR; and NM_000901 for human MR.

14
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16 **Strong response to 3-keto-steroids by sturgeon and gar mineralocorticoid receptors**

17 We examined steroid-inducible transcriptional activation of gar and sturgeon MRs
 18 using the MMTV-driven reporter construct [33, 44]. For comparison, we also examined
 19 transcriptional activation of human MR and zebrafish MR. At 1 nM, Aldo, B, S, DOC, F and
 20 Prog were strong inducers of luciferase activation by gar MR and sturgeon MR and by zebrafish

1 MR, with the exception of Prog which had a lower signal. These MRs show little stimulation
2 by 1 nM DHT and E2 (Figure 4). At 1 nM, Aldo, B, DOC were strong transcriptional activators
3 of human MR, which was activated to a lesser extent by S, and weakly activated by Prog (Figure
4 4).
5



6 **Steroid Concentration = 1 nM**

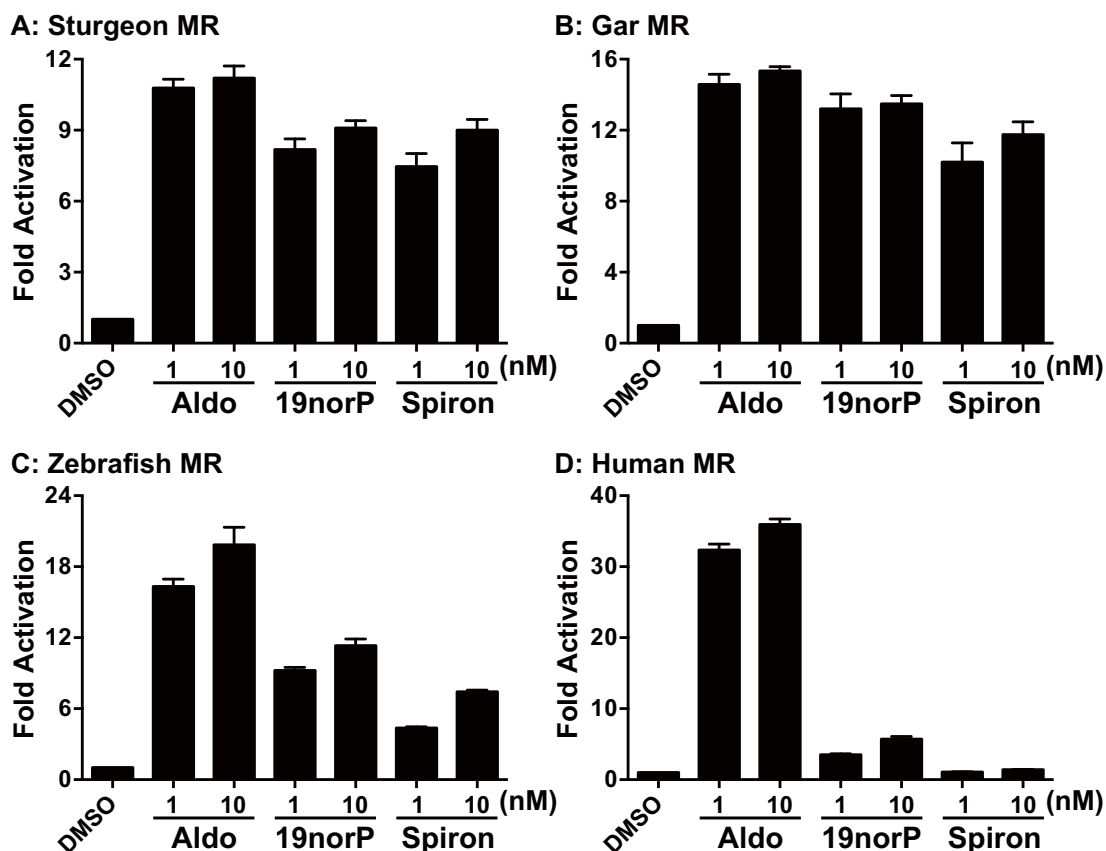
7 **Figure 4. Ligand-specificities of fish and human MRs.**

8 Full-length sturgeon MR (A), gar MR (B), zebrafish (C), and human MR (D) were expressed in
9 HEK293 cells with an MMTV-luciferase reporter. Cells were treated with 10^{-8} M Aldo, F, B, S,
10 DOC, Prog, 5α -dihydrotestosterone (DHT), 17β -estradiol (E2) or vehicle alone (DMSO).
11 Results are expressed as means \pm SEM, n=3. Y-axis indicates fold-activation compared to the
12 activity of control vector with vehicle (DMSO) alone as 1.
13

14 The agonist activity of Prog for sturgeon, gar and zebrafish MRs and for trout MR [16]
15 and the evidence that spiron is an agonist for trout [16] and zebrafish MRs [19] stimulated us to
16 examine transcriptional activation of sturgeon and gar MRs by spiron, an agonist for human MR.
17 We also investigated transcriptional of sturgeon, gar and zebrafish MRs by 19norP, which, like
18 Prog and spiron, is an agonist for human Ser810Leu-MR [27]. As shown in Figure 5, spiron

1 and 19norP are agonists for sturgeon, gar and zebrafish MRs.

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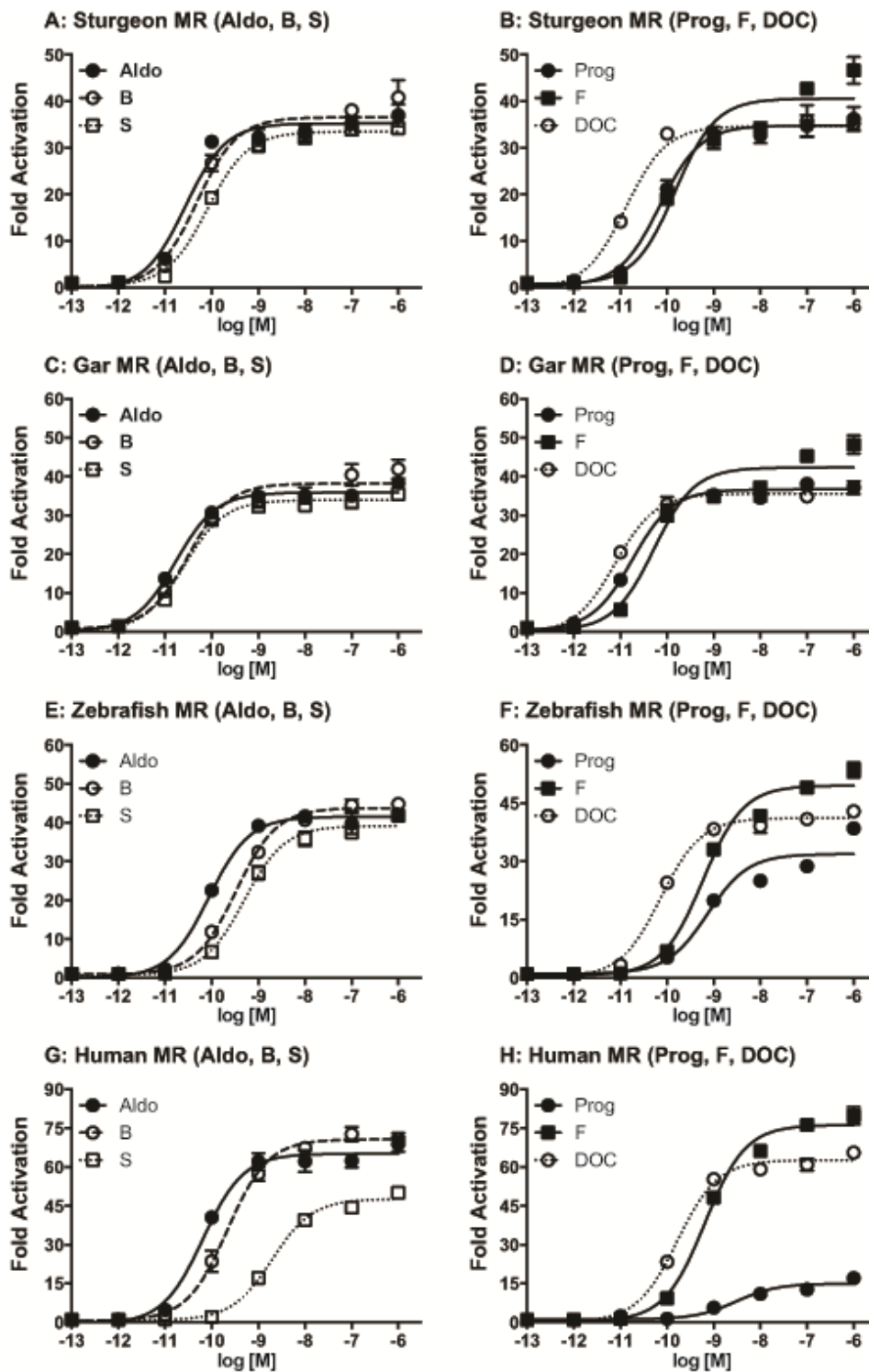
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5 **Figure 5. Transcriptional activation by 19nor-progesterone and spironolactone of sturgeon,**
6 **gar, zebrafish and human MRs.**

7 Full-length sturgeon MR (A), gar MR (B), zebrafish MR (C), and human MR (D) were
8 expressed in HEK293 cells with an MMTV-luciferase reporter. Cells were treated with 1 nM or
9 10 nM Aldo, 19norP or spiron or vehicle alone (DMSO). Results are expressed as means \pm
10 SEM, N=3 and represent fold-activation compared to the control vector with vehicle.

11

12 We examined concentration-dependent activation of gar, sturgeon, zebrafish, and
13 human MRs by Aldo, F, B, DOC, S and Prog (Figure 6, Table 1). Both gar and sturgeon MRs
14 had similar low EC50s, which varied from 7.7 pM to 150 pM for these steroids. For each
15 steroid, the EC50s for gar MR were a little lower than for sturgeon MR.



1

2 **Figure 6. Concentration-dependent transcriptional activities of fish and human MRs.**

3 Concentration-response profiles of full-length sturgeon MR (A and B), gar MR (C and D),
4 zebrafish MR (E and F), and human MR (G and H) for various steroids. HEK293 cells were
5 transiently transfected with the MMTV-containing vector together with an MR expression vector.
6 Cells were incubated with increasing concentrations of Aldo, B, and S (A, C, E, and G) or Prog,

1 F, and DOC (B, D, F, and H) (10^{-13} to 10^{-6} M). Data are expressed as a ration of steroid to
 2 vehicle (DMSO). Each column represents the mean of triplicate determinations, and vertical
 3 bars represent the mean \pm SEM.

4
 5 **Table 1. EC50 activities for 3-keto-steroid transcriptional activation of sturgeon, gar,**
 6 **zebrafish and human MRs**

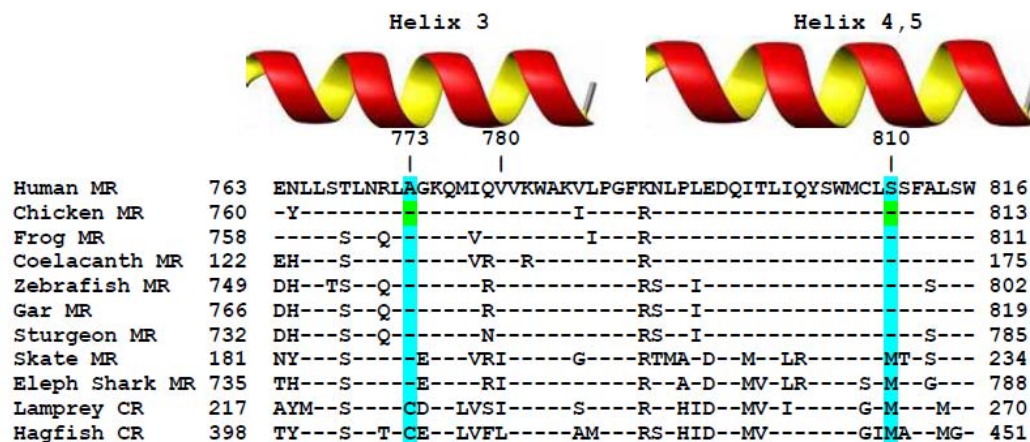
	Aldo	DOC	B	S	F	Prog
Sturgeon MR	2.7×10^{-11}	1.3×10^{-11}	4.8×10^{-11}	8.2×10^{-11}	1.5×10^{-10}	7.0×10^{-11}
Gar MR	1.7×10^{-11}	7.7×10^{-12}	3.1×10^{-11}	2.6×10^{-11}	5.3×10^{-11}	1.8×10^{-11}
Zebrafish MR	8.8×10^{-11}	7.4×10^{-11}	3.3×10^{-10}	5.0×10^{-10}	5.9×10^{-10}	7.4×10^{-10}
Human MR	6.5×10^{-11}	1.7×10^{-10}	2.2×10^{-10}	2.0×10^{-9}	6.5×10^{-10}	-

7

8

9 In comparison, EC50s of Aldo, B and F were similar for zebrafish and human MR and
 10 a little higher than their EC50s for sturgeon and gar MR. EC50s of DOC, S and Prog for
 11 zebrafish MR were higher than their EC50s for sturgeon and gar MR, but lower than the EC50s
 12 for human MR. Prog had a lower, but still significant, maximal activation for zebrafish MR
 13 while 100 nM Prog had little activation of human MR. Overall all corticosteroids and Prog had
 14 EC50s that would be consistent with a physiological role in transcription of the MR in sturgeon,
 15 gar and zebrafish (Table 1, Figure 6).

16 In human MR, Ser-810 and Ala-773 are important in the low transcriptional activity of
 17 Prog. Prog, spiron and 19norP can activate human MR with selective mutations at either
 18 Ser-810 or Ala-773 [27, 37, 38]. For example, at 1 nM, prog, spiron and 19norP are agonist for
 19 a Ser810Leu mutant MR [27, 37, 38]. We extracted the sequence of helices 3-5, which contain
 20 Ser-810 and Ala-773, from sturgeon, gar and zebrafish MR (Figure 7) and other teleosts. All of
 21 these ray-finned fish contain a serine and alanine that aligns with Ser-810 and Ala-773 in human
 22 MR.



23

24 **Figure 7. Alignment of vertebrate MRs to Serine-810 and Alanine-773 in**

1 **human MR.**

2 Human Ser-810 and Ala-773 are conserved in ray-finned fish MRs. Skate MR, elephant shark
3 MR, lamprey CR and hagfish CR contain a methionine corresponding to human Ser-810.
4 Lamprey CR and hagfish CR contain a cysteine corresponding to Ala-773 in human MR.
5 Amino acids that are identical to amino acids in human MR are denoted by (-).
6

7 **DISCUSSION**

8 Several corticosteroids are physiological activators of the MR in cartilaginous fishes,
9 ray-finned fishes and tetrapods. Interestingly, Aldo, the mineralocorticoid for terrestrial
10 vertebrates, first appears in lungfish [45]. Nevertheless, Aldo is a potent activator of the
11 lamprey CR [6], which is ancestral to the MR [5-7, 11]. Interestingly, F, DOC, B and S and
12 Prog also are transcriptional activators of the CR in lamprey and hagfish [6], with only S, thus far,
13 found to have mineralocorticoid activity in lamprey [12, 13]. In skate, which has separate MR
14 and GR genes, Aldo, F, DOC and B are strong transcriptional activators of the MR [14]. F,
15 DOC, B, S and Prog are found in teleosts [26] and F and DOC have been proposed to be
16 transcriptional activators of teleost MRs [4, 15-19, 21, 22, 24, 25].

17 Absent, until now, was information about the response to corticosteroids of MRs in
18 sturgeon and gar, two basal fish that fill in the gap between elasmobranchs and teleosts (Figure 2).
19 Here we report that sturgeon MR and gar MR have EC₅₀s below 1 nM for Aldo, F, DOC, B, S
20 and Prog. Interestingly, we find that zebrafish MR also has a similar strong response to these
21 corticosteroids and Prog. Moreover, spiron and 19norP are agonists for sturgeon, gar and
22 zebrafish MRs. This low selectivity for 3-keto-steroids (Figure 1) that can activate these fish
23 MRs resembles the response to these steroids by lamprey and hagfish CR [6] and skate MR [14].
24 Thus, this strong response of the MR to a broad panel of 3-keto-steroids was conserved after the
25 third whole-genome duplication at the base of the teleosts [30-32, 46].

26 In contrast, human MR is more selective for 3-keto-steroids with higher EC₅₀s for S and
27 Prog. Our data showing weak activation by Prog and 19norP of human MR is in agreement
28 with other studies [35-37]. The strong response to Prog, spiron and 19norP of ray-finned fish
29 MR is interesting in the light of the report by Geller et al. [27] that human MR with a Ser810Leu
30 mutation was activated by 1 nM Prog, 19norP and spiron. Mutagenesis studies and structural
31 analyses of the MR-Leu810 mutant led to the hypothesis that Leu-810 on α -helix 5 has
32 stabilizing van der Waals contacts with Ala-773 on α -helix 3 [27, 37, 38] to explain the strong
33 transcriptional activation by Prog, 19norP and spiron. This serine and alanine are conserved in
34 and sturgeon and gar MRs, as well as in zebrafish MR (Figure 7) [4] and other teleost MRs
35 indicating that other mechanism(s) can lead to a strong response of sturgeon, gar and zebrafish
36 MR to Prog, 19norP and spiron. Activation by Prog of zebrafish MR is of concern because

1 zebrafish is an established model system for studying gene regulation in teleosts, as well as
2 providing insights into human physiology [47]. Prog activation of zebrafish MR may confound
3 data that focuses on activation of the PR. Prog may also be an agonist for the MR in medaka
4 and other teleosts that have a serine and alanine that correspond to Ser-810 and Ala-773 in
5 human MR.

6

7 **Mechanisms for regulation of steroid activation of ray-finned fish MR**

8 The strong response of zebrafish MR, as well as sturgeon and gar MRs, to five
9 corticosteroids, Prog, 19norP and spiron requires one or more mechanisms to provide
10 steroid-specific regulation of transcriptional activation of these ray-finned fish MRs. At this
11 time, such mechanisms in gar, sturgeon and zebrafish MRs or other ray-finned fish MRs are
12 poorly understood. Clues for possible mechanisms may be found from insights into regulation
13 of mammalian MRs [4, 11, 48-53]. One possibility is an important mechanism in epithelial
14 cells for regulating access of F and B to mammalian MR by tissue specific expression of
15 11 β -HSD2, which selectively converts F and B, respectively, to cortisone (E) and
16 11-dehydrocortisone (A), two inactive steroids. Aldo is inert to 11 β -HSD2, allowing Aldo to
17 occupy the MR in epithelial cells in which 11 β -HSD2 inactivates F and B [11, 51, 52].
18 11 β -HSD2 is found in ray-finned fish [54, 55], including sturgeon (Accession: ALE30175) and gar
19 (Accession: XP_006641583). Expression of 11 β -HSD2 in MR-containing tissues provides a
20 mechanism to exclude F and B from the MR. DOC, S and Prog, which have low EC₅₀s in gar,
21 sturgeon and zebrafish, lack an 11 β -hydroxyl group and are inert to 11 β -HSD2.

22 Other regulatory mechanisms of the response of the MR to 3-keto-steroids include
23 tissue-selective synthesis of 3-keto-steroids [5, 56-58], selective sequestration of 3-keto-steroids
24 to plasma proteins [48, 53, 59], steroid-specific conformational changes that regulate MR
25 binding of co-activators [60-64], effects of inter-domain interactions between the NTD and the
26 LBD [19, 50, 64, 65] and post-translational modifications, such as phosphorylation, and
27 SUMOylation [49, 66, 67].

28

29 **Authors Contributions**

30 A.S., K.O., R.S., and S.A. carried out the research. M.E.B. and Y.K. conceived and
31 designed the experiments and wrote the paper. All authors gave final approval for
32 publication. We have no competing interests.

33

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7

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