Differential expression of two glucocorticoid receptors in seabass (teleost fish) head kidney after exogeneous cortisol inoculation

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A B S T R A C T

Stressful conditions include a prompt release of corticosteroid hormones which can mediate gene expression through glucocorticoid receptors (GR). Since two seabass (Dicentrarchus labrax) GRs have been cloned and sequenced from peritoneal cavity cells (DlGR1) and liver (DlGR2), a comparative amino acid sequence analysis that included Haplochromis burtoni HbGRs, was carried out and homologies disclosed. The DlGR1 and DlGR2 deduced aminoacid sequences showed 61% identity (I) and 70% similarity (S). Moreover, DlGR2 was similar to HbGR2b (69% I, 73% S), and the DlGR1 to HbGR1 (72% I, 78% S). In addition, we examined the expression of the DlGRs after exogeneous cortisol inoculation into the peritoneal cavity, mimicking stress effects. At various times after the administration (3 h, 24 h, 1 week), gene expressions was evaluated in head kidney by real-time PCR. In addition, immunoblotting and densitometry analyses were performed with anti-DlGR1 antibodies. Although sea bass head kidney expressed both DlGR1 and DlGR2 they were differentially modulated by intraperitoneal implant of exogeneous cortisol.

1. Introduction

Glucocorticoids (GCs), including cortisol and corticosterone, are secreted by the cortex of the adrenal gland and exert a pivotal control role in vertebrate physiology. GCs and the nuclear glucocorticoid receptors (GRs) are involved in the regulation of cell growth, bone density, metabolism and cardiovascular system, and influence behavior (Charmandari et al., 2005). In addition, elevated GCs levels exert immunosuppressive effects (Wendelaar Bonga, 1997; Mommsen et al., 1999; Sapolsky et al., 2000). Moreover, it is a key mediator of stress-associated responses (Vijayan et al., 1999; Prunet et al., 2006; Bury and Sturm, 2007).

A GR phylogenetic tree, distinguishes a cluster of tetrapod GRs and a cluster of the teleost fish GRs (Stolte et al., 2006). The fish GR cluster is subdivided into two different clades representing two different GR genes. Duplicate GR genes have been found in salmonids (Bury et al., 2003; Ducouret et al., 1995) and cichlids including Burtons’ mouthbrooder (HbGRs) (Greenwood et al., 2003). GRs display high aminoacid sequence identity especially in the C-terminal part. Finally, tissue GR analyses revealed differential distribution and expression levels of duplicate genes (Bury et al., 2003; Greenwood et al., 2003; Ducouret et al., 1995).

In seabass (Dicentrarchus labrax), two distinct GRs have been sequenced: the DlGR1 from the peritoneal cavity cells (Vazzini et al., 2005) and a GR (here named DlGR2) from the liver (Terova et al., 2005). DlGR1 cDNA disclosed four functional domains, for transcriptional activation, DNA-binding, nuclear localization, and hormone-binding; it is expressed in macrophages and neutrophils from the peritoneal cavity (Vazzini et al., 2007), in the head kidney, spleen, gill, intestine, heart and liver tissues (Vazzana et al., 2008). Although stress-associated increased plasma cortisol can modify the GR gene expression, contrasting results on gene regulation have been reported (Terova et al., 2005; Vijayan et al., 2003; Sathiya and Vijayan, 2003; Takahashi et al., 2006). In the present paper, DlGR1, DlGR2 and HbGR sequences were compared and homology levels disclosed. Furthermore, since the differential expression of the duplicate GR genes suggests that they could regulate different functions, the DlGR1 and...
DlGR2 expressions following a cortisol stimulus were examined. Our results show that the head kidney, the main fish hemopoietic organ involved in immune responses, expresses both DlGR1 and DlGR2 differentially modulated by intraperitoneal implant of exogenous cortisol that mimics stress effects.

2. Materials and methods

2.1. Animals and experimental design

Sea bass (D. labrax, Perciformes, Moronidae) (200–250 g) were provided by ECO-ITTICA (Marsala, TP) fish farm. Three distinct fish groups were arranged in separated 100 L tanks with continuously flowing aerated sea water (18 °C) and acclimated for at least two weeks before starting the experiments. Each group contained 15 individuals/experiment for each tank, and a total of 45 fish was examined. The groups were designed as untreated (Untr.), vehicle-treated (I) and cortisol-treated (II).

To avoid circadian effects, fish sampling was carried out at a set up time in the morning. Fish were anaesthetized with 0.05% 3-amino benzoic acid ethyl ester (Sigma-Aldrich, Germany) in seawater. Intraperitoneal implants of 20 mg cortisol kg$^{-1}$ body mass in 2.5 mL medium (ethanol/1:4 diluted Hanks balanced salt solution, v/v) (vehicle) were given to each specimen from group II. Fish from group I (control) were treated with the vehicle alone (sham fish). Five fish from I and II groups were sampled at 3 h, 24 h and 1 week after the inoculation, and five fish were taken at the same time/point from the untreated group (Untr.). After anesthesia, bleeding fish were killed with a lethal dose of 0.1% 3-amino benzoic acid ethyl ester, and the head kidney was excised. To prepare plasma, the blood was withdrawn from the heart into heparinized sterile plastic syringes, and centrifuged at 800×g for 10 min at 4 °C. The excised head kidney was divided in two parts, one half was placed in RNA later (Ambion) for mRNA analysis, and the second part was stored at −80 °C for protein analysis.

2.2. Plasma analysis

Plasma cortisol was measured by using the ELISA method according to the manufacturer instructions (Globe diagnostics Srl, Italy). This test is based on competition enzyme immunoassay into microwells coated with specific anti-cortisol-antibodies simultaneously with conjugated cortisol-peroxidase. After washing, the remaining enzymatic activity bound to the microwell surface was detected and quantified by adding a chromogen–substrate mixture and stop solution. Optical density was recorded at 450 nm, and cortisol concentration was calculated based on a series of standards.

The glucose and lactate levels were recorded by Accutrend GC kit (Boehringer Mannheim) and Lactate Dry-Fast kit (Sentinel diagnostics, USA). The glucose and lactate levels were recorded by Accutrend GC kit (Boehringer Mannheim) and Lactate Dry-Fast kit (Sentinel diagnostics, USA). The glucose and lactate levels were recorded by Accutrend GC kit (Boehringer Mannheim) and Lactate Dry-Fast kit (Sentinel diagnostics, USA).

2.3. Total RNA extraction and cDNA synthesis

Total RNA was isolated from the head kidney, by using a RNAqueous™-Midi Kit purification system (Ambion), and reverse-transcribed by the Kit Ready to Go™-T-primed first-strand using random primers (Amersham-Pharmacia Biotech, USA).

2.4. Real-time PCR analysis

Tissue expression of DIGR genes (DlGR1 Accession Number AY619996; DlGR2 (liver GR) Accession Number AY549005) was detected by real-time PCR by using the Taqman method. Primers and hybridization probes were designed using primer express software V.0 and synthesized commercially (Applied Biosystem, Foster City, CA, USA). The Taqman probe sequence contained a 5′-FAM fluorophore and 3′ MGB quencher for target genes (DlGR1 and DlGR2) and 5′-VIC fluorophore and 3′ MGB quencher for the housekeeping gene (D. labrax actin Accession Number AY148350). Real-time PCR analysis was performed using the Applied Biosystem 7500 real-time PCR system. Tissue expression was performed in a 25 µL PCR reaction containing 2 µL cDNA converted from 250 ng of total RNA, 100 nM of DIGR1 probe (TGGTCTGGTGGCGGTCT), 400 nM of DIGR1 forward (5′-TGGGGCCCAAAG-3′) and reverse primers (5′-TCTGA-TACTGGCTTACAAAGG-3′), 100 nM DIGR2 probe (CCACCGCTTCC), 400 nM of DIGR2 forward (5′-CAACAGAGGCGGGTAGTAC-3′) and reverse primers (5′-CCCCAGGCTGCTACAAAGA-3′), 100 nM of actin probe (ACCAACGGCAGAGG), 400 nM of actin forward (5′-CAGAGGCTGGCTACTCTTCA-3′) and reverse primers (5′-TCCTGTGATCTCAGGAGAT-3′), and 12.5 µL of Taqman PCR Master Mix (Applied Biosystem). The 50 cycles of the two steps PCR program consisted of initial polymerase activation for 3 min at 95 °C followed by denaturing at 95 °C for 15 s, and annealing/extension at 60 °C for 45 s, in which fluorescent signal was detected.

For every time point/treatment five specimens were used and each cDNA sample was run in triplicate together with negative controls. To obtain sample quantification the RQ = 2$^{-\Delta\Delta C_{t}}$ method was used and the relative changes in gene expression analyzed as described in the Applied Biosystem Use Bulletin N.2 (P/N 4303859). For the Ct calculation to be valid, the amplification efficiencies of the target and the reference gene must be approximately equal. To assess whether amplicons have the same efficiency, a sensitive method was used for checking how Ct varies with template dilution. Serial dilutions (1, 0.5, 0.2, 0.1, 0.005, 0.002, 0.001) of cDNA were amplified by real-time PCR using the target and the housekeeping gene specific primers and the Ct, i.e., $C_{t,DlGR1,actin} = C_{t,DlGR2,actin}$, was calculated, in 3 replicates, for each cDNA dilution. Data were analyzed using the linear regression analysis. The amount of DIGR1 and DIGR2 transcripts was normalized with actin to compensate for variations in the amount of RNA. Relative DIGR1 and DIGR2 expressions were determined by dividing the normalized value of DIGR1 and DIGR2 by the normalized value obtained from the untreated tissue.

2.5. Sequence comparison

A BLAST search was performed for comparing sequences. Multiple alignments were accomplished with the Clustal W program (Thompson et al., 1994), and similarity and identity percent values were recorded.

2.6. Tissue homogenate supernatant (THS) preparation

Tissue samples were crushed in liquid nitrogen, and kept in ice for 1 h with 1 mL of lysis buffer (RIPA: 0.5% sodium deoxycholate minimum 97%; 1% NP40; 0.1% SDS with PBS-T (Na$_2$HPO$_4$ 1 M, NaH$_2$PO$_4$ 1 M, NaCl 1.5 M, 0.1% Tween 20) supplemented with a protease inhibitor cocktail (pepsatin A, E-64, bestatin, leupeptin, aprotinin, and AEBSF) diluted 1:200 (lysis buffer)). Then the samples were centrifuged at 15,000×g for 30 min at 4 °C. The supernatants were collected, dialyzed against 50 mM Trizma base (Tr[hydroxymethyl] aminomethane) Sigma-Aldrich, pH 7.5, and the total protein content was determined by using the method of Bradford (1976).

2.7. SDS-PAGE and Western blot

SDS-PAGE (7.5%) under reducing conditions was performed according to the method of Laemmli (1970), and samples, adjusted to 25 µg protein content, were analyzed. The protein pattern of THS was transferred to a nitrocellulose membrane by using a semidyrid transfer apparatus (BioRad) and blocked with 5% bovine serum.
albumin (BSA, Sigma-Aldrich) in TBS-T (20 mM Trizma base pH 7.5, 300 mM NaCl, 0.1% (v/v) Tween 20 with 0.02% sodium azide) for 1 h at room temperature (r.t.).

To identify DIGR1, the membrane was incubated with a specific anti-DIGR1 antiserum (1:3000 dilution, 3 h at r.t.), washed with TBS-T (three times for 5 min), incubated with alkaline phosphatase-conjugated goat anti-rabbit IgG (Sigma-Aldrich, 1:10000 for 1 h at r.t.), and washed with TBS-T (three times for 5 min). The membrane was incubated with 5-bromo-4-chloro-3-indolyl phosphate/nitro blue tetrazolium liquid substrate system (BCIP/NBT Sigma-Aldrich). To check for antibody specificity, the anti-DIGR1 antiserum was substituted with preimmune rabbit IgG at a suitable dilution. To check for unspecific cross-reaction of the secondary antibody, the primary antibody was not added.

The specific anti-DIGR1 antibody had been previously prepared and already used for DIGR1 immunohistochemical identification studies as reported by Vazzana et al. (2008). In brief, anti-DIGR1 polyclonal antibody was raised in rabbit by using a peptide as antigen corresponding to the hydrophilic sequence located in the N-terminal region (transcriptional activation domain) of DIGR1 (residues from 85 to 98: LEDHESRLTRDQK). The peptide sequence was selected (by antigen-prediction program) and synthesized by Sigma Genosys (UK).

The integrated density value (IDV) of immunoblots was determined using AlphaImager™ software.

2.8. Statistical analysis

The experiments were performed using 5 specimens for each time point and the results were expressed as a mean value ± SD. The data were compared using the analysis of variance (ANOVA). Homogeneity of variances was performed using Cochran’s test prior to the ANOVA analysis. Whenever necessary the data were log-transformed. Post-hoc comparisons were made using Student’s t test and the significant level was set at p<0.05; the analysis was performed with Statistica 6.0 (StatSoft, Tulsa, OK, USA). Linkage between plasma cortisol levels and gene expression was analyzed at 3 h, 24 h and 1 week p.i. in five specimens from each group. The cortisol treatment enhanced the expression of both DIGR1 and DIGR2 genes that reached the highest levels at 24 h (p<0.001, p<0.01) (Fig. 2A, B). While the DIGR1 gene expression (Fig. 2A) increased significantly at 24 h and at 1 week, the DIGR2 gene expression (Fig. 2B) increased significantly (p<0.05) at 3 h and reached the untreated values at 1 week. Until the end of the experimentation, sham fish presented no significant variation in the expression of DIGR1 gene (Fig. 2A), while a significant (p<0.05) increase in the DIGR2 gene expression was found in the head kidney from sham fish at 3 h and 1 week p.i (Fig. 2B).

Moreover, significant differences (p<0.01) between the expressions of DIGR1 and DIGR2 genes were found at 24 h and 1 week post inoculation with cortisol.

Statistical analysis disclosed that there was a correlation (R² = 0.49) between enhanced DIGR1 gene expression and high plasma cortisol levels recorded at 24 h (Figs. 1A and 2A). Such a correlation was not found between the cortisol level and the DIGR2 gene expression.

3.4. DIGR1 protein abundance in the head kidney tissue

Since anti-DIGR1 specific antibodies were available, an immunoblotting analysis was carried out. As shown in Fig. 3, the head kidney homogenate preparations from sham fish (group I) presented a significantly lowered band density for each time points. The lowest IDV were found in the control samples (group I) both at 24 h and 1 week (p<0.001) (Fig. 3).

4. Discussion

As predicted by previous reports (Greenwood et al., 2003; Stolte et al., 2006) on GR molecular phylogenesis, teleost and nonteleost GRs form distinct branches. GR gene duplicate copies (DIGR1 and DIGR2) that have been found in sea bass (Vizzini et al., 2007; Terova et al., 2005) are in accordance to the GR gene duplication within the teleost lineage. We compared the deduced amino acid sequences of D. labrax (Perciformes, Moronidae) DIGR1 (Vizzini et al., 2007) and DIGR2 (indicated as GR by Terova et al., 2005), and both were compared to HbGR1, HbGR2a and HbGR2b isoforms of H. burtoni (Perciformes, Cichilidae) (Greenwood et al., 2003).

As shown by an alignment of the predicted amino acid sequences, the DIGR duplicate copies have highly conserved aminoacid sequences in C-region, in particular the DNA-binding domain and
the hormone-binding domain are the best conserved (84% I, 85% S and 88% I, 95% S), whereas a lower identity is shared by the nuclear localization domains (47% I and 62% S). In N-region the transcriptional activation domain of the two sequences presents the highest diversity (29% I and 45% S). Nuclear localization and transcriptional activation domain differences could be related to a differential expression of the duplicate genes regulated by the same hormone. A similar conclusion has been reported by Greenwood et al. (2003) on the domains of HbGR1 and HbGR2.

Compared to other fish species (Stolte et al., 2006), DlGR2 gene shows a 27-nucleotide insert that encodes nine additional amino acids (WRARQNTDG). This sequence is remarkably conserved among teleostean fish (Stolte et al., 2006). Although Stolte et al. (2006) proposed to name GR1 the gene containing the insert and GR2 the duplicate form, to avoid confusion with our previous papers we named “DlGR1” (Vizzini et al., 2007) the GR that does not present the insert, and “DlGR2” the GR (Terova et al., 2005) that contains the insert.

The DlGR1, DlGR2, HbGR1 and HbGR2 amino acid sequence alignments show that the DlGR1 is more similar to the HbGR1 (72% I–78% S) than to the DlGR2 (70% I–61% S). The DlGR2 is similar to the HbGR2, mainly to the isoform b that contains the insert.

According to Stolte et al. (2006), differences in receptor protein function could be indicated by differential expression of the duplicate DlGR genes.

It is well documented that in fish, cortisol is the major stress-related hormone, and treatment with exogenous cortisol mimics stress effects and exerts, in distinct species, differential GR genes regulation (Vijayan et al., 2003; Sathiaya and Vijayan, 2003; Takahashi et al., 2006). In cortisol-treated sea bass, plasmatic parameters supported that stress conditions had been reached. According to previous papers on fish population density, glucose, lactate and osmolarity levels were enhanced (Mommsen et al., 1999; Milligan and Girard, 1993; Vazzana et al., 2002). High values, including endogeneous cortisol level, found in the plasma of sham fish indicated that the inoculation procedure caused a moderate stressing effect as compared to the lower levels recorded in the plasma of the untreated fish. The further increase in stress indicators levels, due to the exogeneous cortisol, disclosed the effects of the hormone. It is noteworthy that no variations were observed in the untreated fish even when assayed at different time points of the experimental schedule. However the average cortisol level (80.5 ng mL⁻¹) of the untreated specimens examined in the present study was higher than that (10±8 ng mL⁻¹) previously reported by Vazzana et al. (2002). Such a difference could be due to different environmental conditions and population density between fish from distinct farms. Furthermore, in the previous study, control fish were maintained in 400 m³ farm tanks whereas, in the present study, 0.5 m³ aquarium tanks were used, therefore environmental conditions cannot be straightly compared. The question arises whether the enhanced plasma cortisol differentially modifies the expression of DlGR genes. In rainbow trout, concentrations of cortisol required to induce activation of downstream genes is significantly different between the duplicate

![Plasma cortisol (A), glucose (B), lactate (C) and osmolarity (D) levels of untreated (Untr.), vehicle-treated (I) and cortisol-treated (II) sea bass at 3 h, 24 h and 1 week. Values represent means±SD (n = 5). The average value of the untreated samples was obtained by putting together results from five untreated specimens at every time point (n = 15). Statistical significant differences between cortisol-treated and vehicle-treated fish are shown upon the bars (*p<0.05, **p<0.01, ***p<0.001). Significant differences between untreated and treated fish (I, II) are also indicated.](image-url)
plasma cortisol displaying a time-course profile. However, although both the DIGR-mRNAs were significantly increased at 24 h p.i., the DIGR1 gene expression was mainly enhanced reaching the highest level. In addition, when (at one week) the expression of both GR genes decreased, the DIGR1 gene expression maintained a high level. The correlation analysis supports that plasma cortisol level and head kidney DIGR1 gene expression are correlated. Since macrophages and neutrophils, components of the head kidney white pulp (Vazzana et al., 2008) and peritoneal cavity, express DIGR1 (Vizzini et al., 2007), a role of this GR on cortisol immunomodulation may be suggested. A similar result has been reported on the gene expression of two functionally distinct rainbow trout GR genes: RtGR2b gene requires a lower level of cortisol than the RtGR1a gene to induce transactivation (Bury et al., 2003).

It is of interest that the DIGR1 protein, identified in the head kidney homogenate supernatant by specific antibody and analyzed by densitometry, appears to be mainly lowered in sham fish at 24 h and at 1 week after the vehicle inoculation. According to Wallace and Cidlowski (2001), the possibility exists that the injected medium (vehicle) does not activate the DIGR1 gene transcription whereas the GR protein could be diminished as an effect of a proteasoma-mediated degradation due to the physiologically enhanced plasma cortisol (250–300 ng ml⁻¹ plasma cortisol). Such a DIGR1 protein diminution could be compensated by the transcription activity enhanced by exceeding plasma cortisol. This is supported by the correlation between plasma hormone level and real-time PCR analysis. In the cortisol-treated specimens, the production of DIGR1 protein appears to be increased but a constant level was found at any time point, like that of untreated specimens.

Our results allow us to speculate that a differential expression of the two DIGR genes could reflect the regulation of tissue specific genes by different types of glucocorticoid-sensitive promoters. In addition, according to the competitive effect of a GR blocker (RU486) on the activity of cortisol-treated peritoneal phagocytes (Vizzini et al., 2007), an immunomodulatory role of the differently expressed head kidney DIGR1 may be suggested.

References


