Retinoic acid regulates the human methionine sulfoxide reductase A (MSRA) gene via two distinct promoters

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ABSTRACT

MSRAs (methionine sulfoxide reductases A) are enzymes that reverse the effects of oxidative damage by reducing methionine sulfoxide back to methionine and recovering protein function. In this study we demonstrate that the transcriptional regulation of the human MSRA gene is complex and driven by two distinct promoters. Both promoters demonstrate high expression in human brain and kidney tissues. The upstream (promoter 1) regulates the msrA1 transcript that codes for the mitochondrial form of MSRA and is highly active in a broad range of cell lines. The downstream promoter (promoter 2) regulates the msrA2/3 transcripts that code for the cytosolic/nuclear forms of MSRA and is generally less active. Promoter 2 contains a 65 bp putative enhancer region that is very active in the retinal pigment epithelium-derived D407 cell line. Both promoters are partially regulated by all-trans retinoic acid via RARA and other RARs.

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Introduction

Methionine sulfoxide reductases (MSRs) are a family of antioxidant enzymes that convert free or protein bound methionine sulfoxide (MetO) back to methionine [1,2]. This process is known to play a critical role in recovering protein functionality and in protection against oxidative stress [3]. There are two distinct subfamilies of MSRs. MSRA are capable of reducing the S diastereomer, Met(S)O while MSRBs reduce the R diastereomer, Met(R)O [4–7].

The importance of MSRs in protection from oxidative stress and in the aging process has been well documented [1,2]. MSRA overexpression increases lifespan and fertility as well as their resistance to oxidative stress and hypoxia. In rats, MSRA levels have been shown to decrease with age [13]. MSRA knockout mice suffer from neurological abnormalities, are more susceptible to oxidative stress and have a 40% reduction in their lifespans [14]. In Drosophila, MSRA overexpression increases lifespan and fertility as well as their resistance to the insecticide paraquat [15]. In human WI-38 fibroblasts, MSRA was found to be downregulated during replicative senescence [9].

Our previous study demonstrated that the MSRA gene contained two putative regulatory regions (promoters) 40 kbp apart which generate three different transcripts [16]. These transcripts generate different protein isoforms differing in their N-termini which in turn determines their intracellular localizations [16]. The main transcript, msrA1 (AY958429), is generated by promoter 1 and codes for the main isoform of MSRA which localizes to the mitochondria [16,17]. The other two transcripts, msrA2 (AY958430) and msrA3 (AY958431), are generated by promoter 2 and code for two isoforms of MSRA that localize to the cytosol and cytosol/nucleus, respectively [16]. The msrA3 transcript was subsequently reported by another group which also determined its nuclear/cytosolic localization [18]. More recently, alternatively spliced forms of mitochondrial msrA have also been identified in the rat [19].

In the retina, MSRA localizes to the retinal pigment epithelium (RPE), photoreceptor synapses and ganglion cells [16] and may be playing an important role in protecting these tissues from oxidative and photo-damage [16,20]. In cultured RPE cells, siRNA-mediated gene silencing increased their susceptibility to tertiary butyl-hydroperoxide [16] and hydrogen peroxide [20] induced cytotoxicity. In the monkey retina, the macular RPE has very high levels of MSRA expression [16]. This suggests that RPE may be an appropriate tissue to study MSRA transcriptional regulation.

The upstream human MSRA promoter (promoter 1) was partially characterized recently [21], but little is known about the putative downstream promoter (promoter 2) previously reported [16]. In this study, we have determined that the putative promoter 2 is indeed capable of initiating the transcription process that generates the msrA2/3 transcripts (nuclear and cytosolic MSRA). We have found that both promoters respond vigorously to all-trans retinoic acid (ATRA) and that promoter 2 contains an enhancer region that may explain the high MSRA expression observed in brain and RPE cells [16,20,22].
Results

Expression of msrA transcripts in different human tissues

In order to determine the tissue distribution of the different msrA transcripts, qRT-PCR was performed on RNA from different human tissues (Fig. 1). We were unable to design a primer set that could unequivocally detect msrA1 so a primer set that detects all of the msrAs was used and compared with a set that specifically detects msrA2/3 (Table 1). Hence, Fig. 1A measures the contribution of both promoters while Fig. 1B measures the contribution of promoter 2 specifically. All measurements were normalized to the 18S ribosomal RNA and the neural retina was given to the retina. B, Brain; C, Cerebellum; H, Heart; K, Kidney; Li, liver; Lu, Lung; Pl, placenta; Pr, prostate; R, Neural retina; SI, Small Intestine; SM, Skeletal Muscle; Sp, Spleen; St, Stomach; T, Testis. The tissues were aligned from high to low and retina is marked with a star.

Activity of MSRA promoters in D407, HEK 293 and SH-SY5Y cell lines

To characterize and compare the MSRA upstream (promoter 1) and downstream (promoter 2) promoters, three cell lines were chosen which originated from tissues of known high MSRA expression. The cell lines were D407 (RPE origin), HEK 293 (human embryonic kidney cells) and SH-SY5Y cells (neuronal origin). Promoters were designed to amplify different areas of both promoters and the fragments cloned into pGL4 luciferase reporter vector (Table 1). The constructs were transfected into the different cell lines and luciferase activity was measured (Fig. 2). The sequences for promoter 1 and 2 were submitted to GenBank and the accession numbers are as follows: EU409840 for promoter 1 (−1919 to −1 bp) and EU409841 for promoter 2 (−1979 to −1 bp). The complete structure for the MSRA human gene was previously published (16) but we have included a brief diagram encompassing the promoter region (Fig. 2). Please refer to reference 16 and the above GenBank files for more specific information.

Constructs 1–2 (promoter 1, −1232 to −1, Fig. 3D) and 2–3 (promoter 2, −971 to −1, Fig. 3D) demonstrated the highest activity (Fig. 3A) in D407 cells. In HEK 293 cells, promoter 1 was approximately 10-fold more active than promoter 2 (Fig. 3B). The construct 2–3 demonstrated a 2.7 fold increase in activity over the control vector (pGL4). The construct 1–1 demonstrated slightly greater activity than 1–2 in HEK 293 cells (Fig. 3B). These results suggest the possibility that an upstream suppressor element is present in the 1–1 construct which is active in D407 cells but not in HEK 293 cells. In SH-SY5Y cells the activity for both promoters was overall lower than in D407 and HEK 293 cell lines (Fig. 3C). In these cells, promoter 1 activity was higher than promoter 2 (2–3 fold) but contrasted sharply with the approximately 10-fold higher activity observed in HEK 293 cells. SH-SY5Y cells demonstrated similar activity for the different promoter 1 constructs (1–1, 1–2 and 1–3). By contrast, the promoter 2 constructs behaved similarly in all the cell lines tested, with construct 2–3 having the highest activity. Overall, the data suggest that MSRA promoters may be regulated differentially depending on the tissue or organ.

Identification of a putative enhancer in the 5′-distal region in promoter 2

In D407 cells, the considerable decrease in the promoter 2 activity observed between the constructs 2–3 and 2–4 (Fig. 3A) suggested a potential enhancer in the region between −971 and −499 (Fig. 3D). To localize the region responsible for this transcriptional increase, four constructs of approximately 130–150 bp each
were made and their luciferase activity measured (Fig. 4, hatched bars, Table 2). Only one of the constructs, between the positions −731 to −598, demonstrated high luciferase activity similar to the activity of the original 2–3 construct (Fig. 4). The 133 bp region was further subdivided into three additional overlapping constructs of approximately 65 bp each (Fig. 4, Table 3). The first 65 bp (−731 to −666) did not have any activity, the middle region construct (−693 to −628) demonstrated approximately 50-fold increase over control and the last construct (−666 to −600) showed only a 10-fold increase in activity (Fig. 4). These results suggest that the 65 bp region between −693 and −628 is critical to the high expression observed in D407 cells.

Characterization of the promoter 2 putative enhancer region (−693 to −628)

To further characterize this 65 bp enhancer region and to identify specific response elements responsible for its high activity, five different deletion constructs were made and analyzed (Fig. 5, Table 4). Computer-aided analyses (www.gene-regulation.com/pub/databases.html) suggested that the sequences TACTTGCT (deletion 1, putative Oct-1 and/or Myf-3 element), GTTATTAAA (deletion 2, putative HNF-1), AGATGTTTTA (deletion 3, putative HNF-3), TTTTCCTTTT (deletion 4, putative GATA-1, Oct-1, ICSBP or RARA), and TATTTGTTTTA (deletion 5, putative Oct-1, C/EBPα or HNF-3) may be areas of interest. All of the
constructs demonstrated increased activity over the empty vector control. Constructs with deletions 1, 2 and 4 demonstrated diminished activity while constructs with deletions 3 and 5 showed greatly increased activity when compared to the original (−693 to −628). Interestingly, deletions 3 and 5 contain a TGGTTTA sequence that may be a suppressor element.

Effect of all-trans retinoic acid (ATRA) on the MSRA promoters

The computer-aided analysis of both promoter regions suggested putative retinoic acid response elements (RARA) in both MSRA promoters. Moreover, analysis of the 65 bp promoter 2 enhancer region and the results from the above deletion constructs (deletion 4, Fig. 5) which interfered with a putative RARA site suggested that these elements may be of importance.

To test this assumption D407 cells were transiently transfected with constructs 1–2 and 2–3 and treated with increasing concentrations of ATRA (0–15 μM), 18 h after transfection. The cells were harvested 24 h after ATRA treatment and luciferase activity was measured (Fig. 6A). Since ATRA may be toxic, cellular dehydrogenase activity, a measure of cell viability, was monitored over the same concentration range (Fig. 6B). The lower concentrations of ATRA (1 and 5 μM) roughly doubled the activity of both MSRA promoters (Fig. 6A) with no effect on cell viability (Fig. 6B). The promoter induction was still measurable at 10 μM but cell viability was reduced by approximately 25% (Figs. 6A and B). At 15 μM the loss in cell viability was approximately 40% and this correlates with the drop in activity observed for both promoters.

Expression of RARs in D407 cells

Since both MSRA promoters responded to ATRA we performed immunoblots using specific antibodies to RARA, RARB and RARG to determine if they were expressed in nuclear extracts from D407 cells (Fig. 7). The immunoblots indicate that all three RARs are expressed in D407 cells (Fig. 7). For subsequent experiments we focused on RARA for two particular reasons: RARA seems to be more abundant than the other RARs and the computer-aided analyses indicated RARA-specific elements.

Effect of RARA on the activity of MSRA promoters

To elucidate the effect of retinoic acid receptors on the activity of the MSRA promoters, RARA was overexpressed in D407 cells. One set of D407 cells was co-transfected with the construct 1–2 (promoter 1, −1232 to −1, Fig. 3D) and another set of cells with the construct 2–3 (promoter 2, −971 to −1, Fig. 2D). In addition, both set of cells received the RARA construct and a control plasmid as a mock transfection control. The cells were treated with 1 μM ATRA 18 h after transfection and the luciferase activity was measured 24 h after the treatment. The results from the experiments are shown in Fig. 8.

Without ATRA treatment, neither promoter responded to the RARA overexpression (Figs. 8A and B, no treatment). As shown above (Fig. 6) treatment with 1 μM ATRA increased the activity of both promoters (Fig. 8). This effect was likely due to the constitutive expression of RARs in D407 cells (Fig. 7). The cells co-transfected with the RARA overexpression plasmid demonstrated an increase in activity for both promoters (Figs. 8A and B) with promoter 2 being slightly more responsive.

To confirm the interaction of RARA with the MSRA promoters, D407 cells were co-transfected with the luciferase construct 1–2 or 2–3, and RARA siRNA or a negative control siRNA. The RARA siRNA used suppressed the native RARA mRNA levels by 70% (data not shown). The RARA siRNA attenuated the response of both promoters to ATRA (Fig. 9). The reduction was approximately 20% without ATRA treatment and around 40% after the ATRA treatments.

The data suggest that RARA mediates the responses of both promoters and promoter 2 may be somewhat more responsive to ATRA than promoter 1.

Induction of native msrA transcripts by ATRA

To demonstrate that the transcriptional induction observed with the luciferase reporter constructs was indeed relevant to MSRA expression, msrA transcripts were measured by qRT-PCR in RNA from D407 cells treated with ATRA (Fig. 10). The cells were treated with 0–
15 μM ATRA for 24 h and 48 h, and the msrA transcripts measured using the same primers and conditions as in Fig. 1 (Table 1). One primer set measured all msrAs (Figs. 10A and B) and the other set measured msrA2/3 from promoter 2 (Figs. 10C and D). Cell viability was determined for both experiments (Figs. 10E and F) by assaying for cellular dehydrogenase activity.

After 24 h all msrAs (msrA 1, 2 and 3) transcripts demonstrated a dose-dependent induction (1.7 to 4-fold) over control at the 5 to 15 μM concentrations (Fig. 10A). The msrA2/3 transcripts from promoter 2 seemed to be more responsive with a 2.5–8 fold increase over controls (Fig. 10C). After 48 h all msrAs increased by approximately 30 fold (Fig. 10B) at 15 μM ATRA. The msrA2/3 transcripts increased by approximately 18 fold (Fig. 10D) with 15 μM ATRA. The cell viability (cellular dehydrogenase activity) was reduced by 20–30% with the 10 and 15 μM ATRA concentrations (Figs. 10E and F).

Discussion

In this study we demonstrate that the MSRA gene is regulated by two distinct promoters. We partially characterized both MSRA promoters with emphasis on the downstream, promoter 2. This second MSRA promoter is particularly interesting because it regulates the transcription of two distinct mRNAs that generate MSRA isoforms with distinct N-termini that targets them to the cytosol and the nucleus [16]. These isoforms have been previously described as originating by alternative splicings [18,19] but we clearly demonstrate that their expression is controlled by a second MSRA promoter. We have also established that retinoic acid is involved in the regulation of MSRA expression via RARA and suspect that other RAR isoforms may also be involved.

The transcriptional regulation of MSRA is not well understood. In Saccharomyces cerevisiae calcium phospholipid-binding protein (CPBP), a homologue of elongation factor 1 gamma, has been shown to regulate MSRA gene expression [24]. Also, in yeast, thioredoxins have been shown to be essential for MSRA transcriptional activity [25]. In Drosophila the hormone ecdysone (essential in insect molting) induced MSRA via the EcR-USP (ecdysone receptor-ultraspiracle) complex [26]. EcR and USP belong to the same family of the mammalian retinoic acids, thyroid hormone and vitamin D receptors [27]. Thus, this previous work indirectly supports our findings regarding the interactions of the MSRA promoters with retinoic acid receptors.

In the human tissues analyzed by real time RT-PCR, brain and kidney have much higher levels of msrAs than other tissues (Fig. 1A). This higher expression in kidney and brain was previously described by RNA dot blot analysis [28]. Kidney seems to be favored by promoter 1 which generates the mitochondrial msrA1 transcript [16,17]. Promoter 2 which controls the expression of the cytosolic and nuclear transcripts msrA2/3 is most highly active in brain followed by the kidney and neural retina (Fig. 1B). In the retina, MSRA is known to be highly expressed in only three cell types, RPE, ganglion cells and photoreceptors [16]. Using the same primer sets as in Fig. 1 and monkey RNA from the RPE and choriocapillaris regions of the retina, all msrAs (msrA1, 2 and 3) are expressed 4.5 fold more than in the neural retina (data not shown). The msrA2/3 transcripts from promoter 2 were 13 fold greater than in the monkey neural retina (data not shown). Moreover, the RPE-derived D407 cells demonstrated the highest level of activity for promoter 2 of all the cell lines tested (Fig. 3). This suggests that MSRA may play an important role in protecting the RPE from oxidative as well as retinoic acid damage.

**Table 3**

List of oligonucleotides used to clone and narrow the promoter 2 enhancer region

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<td>−731 to −666</td>
<td>CTAGTACTATGTGACACTGAGTGTCAAAATCTGAGAATATCCGGTTATTTATATCCAAAGACCTGATGGCAGCCTACGATCCTCA</td>
<td>AGGTGTTCAATTTAATAACAGCAAGTATCTTTTCATATTTTGACACCTCAAGTTCCAGCATAGTCA</td>
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<td>−693 to −628</td>
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<td>−666 to −600</td>
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**Fig. 5.** Characterization of promoter 2 enhancer region. (A) Sequence of putative enhancer region marking the deleted regions 1 thru 5. (B) Luciferase activity of the 5 deletion constructs (Table 4). All the results were normalized to the Renilla luciferase internal control and related to pGL4 empty vector. Data are expressed as mean ± SD of three measurements.
The transcriptional regulation of MSRA is complex and there is evidence of tissue-specific suppressors (Fig. 3). In D407 cells there is a doubling in promoter 1 activity between constructs 1–1 and 1–2 (Fig. 3A). This suggests the presence of a suppressor(s) between −1232 and −1919 which affects D407 cells but not HEK 293 and SH-SY5Y (Fig. 3). In HEK 293 cells there was a sharp decrease in activity between 1–2 and 1–3 (−1232 to −408). De Luca et al. [21] also in HEK 293 cells demonstrated little change in activity between −1341 and −309 and a large increase in activity between −309 and −155. Since De Luca et al. [21] did not test any constructs between −1341 and −309 and we did not test any promoter 1 constructs shorter than −408, it is difficult to make any direct comparison with our present results.

Promoter 2 also seems to have some suppressor activity between −971 and −1979. This seems to be a general effect since it is similar in all cell types (Fig. 3). Promoter 2 has only 2 to 3-fold increased activity over control in HEK 293 and SH-SY5Y cells. However, construct 2–3 has approximately 25-fold increased activity over control in D407 cells. The reason(s) for this difference are unclear but cell-specific interactions with the enhancer between −693 and −628 (Fig. 3) may provide a partial explanation.

The 65 bp enhancer region identified in promoter 2 seems to interact with multiple transcription factors. Deletion constructs in the region between constructs 2–3 and 2–4 (Fig. 3) identified an area of approximately 65 bp between −693 and −628 that provided high

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Table 4
List of oligonucleotides used to delete different putative elements within the promoter 2 enhancer

![Fig. 6](image-url) Effect of all-trans retinoic acid (ATRA) in the activity of MSRA promoters. D407 cells were transfected with constructs of promoter 1 (construct 1–2, −1232 to −1) and promoter 2 (construct 2–3, −971 to −1). The cells were treated with ATRA 18 h after transfection and luciferase activity was measured 24 h after ATRA treatment. (A) Relative luciferase activity of promoter 1 and 2 constructs after treatment with ATRA. (B) Cell viability (cellular dehydrogenase activity) of the cells at the different ATRA concentrations. Data (mean ± SD of three measurements) are expressed as the fold of luciferase activity related to the activity of non-treated cells.

![Fig. 7](image-url) Expression of retinoic acid receptors in D407 nuclear extracts. RARA, RARB and RARG were detected by immunoblot using antibodies specific to each subtype. Each lane contains 15 μg of nuclear protein extract. For more details see Materials and methods.

![Fig. 8](image-url) Effect of RARA overexpression on the activity of the MSRA promoters. RARA expression construct was co-transfected with promoter 1 and promoter 2 luciferase reporter constructs (1–2 and 2–3, respectively) in D407 cells. The cells were treated with 1 μM ATRA 18 h after transfection and analyzed 24 h after ATRA treatment. (A) Promoter 1 response. (B) Promoter 2 response. For more details see Materials and methods. Data (mean ± SD of three measurements) are representative of three independent experiments.
levels of expression (approximately 50-fold over controls). Further deletions within the 65 bp enhancer region failed to completely obliterate the activity (Fig. 4) and in two instances (deletions 3 and 5) it was further increased. This region also has putative RARE elements [29] and the deletion that disrupted these elements (Fig. 5) decreased but did not eliminate the activity. This suggests that this putative enhancer region is very important to the function of promoter 2.

ATRA, the natural agonist of RARs [29] has an important effect on the transcriptional regulation of MSRA. D407 cells transfected with promoter 1 and promoter 2 luciferase reporter constructs (1–2 and 2–3, respectively) and a siRNA for RARA or a negative siRNA control. The RARA siRNA knocks down approximately 70% of the native RARA mRNA levels. After the co-transfection (24 h), the cells were treated with 0.5, 1 and 5 μM ATRA and luciferase activity was measured 24 h after the ATRA treatment. (A) Promoter 1 response. (B) Promoter 2 response.

Fig. 9. Effect RARA siRNA knockdown on the activity of MSRA promoters. D407 cells were co-transfected with promoter 1 and promoter 2 luciferase reporter constructs (1–2 and 2–3, respectively) and a siRNA for RARA or a negative siRNA control. The RARA siRNA knocks down approximately 70% of the native RARA mRNA levels. After the co-transfection (24 h), the cells were treated with 0.5, 1 and 5 μM ATRA and luciferase activity was measured 24 h after the ATRA treatment. (A) Promoter 1 response. (B) Promoter 2 response.

ATRA also had an effect on the native MSRA promoters in a dose-dependent manner (Fig. 10). The response for promoter 2 was detectable at 1–5 μM in 24 h and before any significant cytotoxicity was detected (Figs. 10C and E). The increased responses at the 1 and 5 μM ATRA concentrations (2–3 fold) were similar to those observed with the shortened promoter constructs in Fig. 5A (please note scale difference). However, both promoters responded vigorously at 15 μM ATRA (Figs. 10B and D) after 48 h. At this concentration cytotoxicity was significant with approximately 30% cell death (Figs. 10E and F). Thus, the 18 and 30-fold increases in activity observed with 15 μM ATRA in 48 h (Fig. 10, promoter 2 and all promoters, respectively) may also be attributed to cell stress and not exclusively to ATRA. This stress response by both promoters needs to be further investigated.

The findings of retinoic acids involvement in the transcriptional regulation of MSRAs are not only novel but potentially very important. Retinoic acids have been shown to modulate anti-oxidant enzymes in different tissues [32,33]. In the retina, ATRA is generated by the photoreceptors and RPE as a by-product of the visual process [34]. Retinoic acids can induce apoptosis in RPE cells [32] by generating reactive oxygen species [32,35]. The generation of reactive oxygen species seems to be due to a direct effect of retinoic acids on the mitochondria [36,37]. Thus, our data suggests that MSRA may be part of a pro-active protective mechanism that responds to ATRA before reactive oxygen species are formed. This may be of particular importance to the retina and RPE cells in particular because of the relative large amounts of ATRA generated in this tissue [34].

Our findings, when taking in context with what is known about MSRA’s function, suggest that this enzyme plays an important role in the anti-oxidative mechanisms of the RPE and possibly in the pathogenesis of RPE-related diseases like age-related macular degeneration [38]. This disease as well as others may benefit from increased expression of MSRA. Thus, understanding the transcriptional regulation of this gene may provide some potential pharmacological solutions to prevent neuronal damage especially in aging diseases.

Materials and methods

Materials

Human total RNAs were purchased from BD Biosciences (Mountain View, CA). Human genomic DNA was purchased from Clontech (Mountain View, CA). Oligonucleotides were purchased from Integrated DNA Technologies, Inc. (Coralville, IO). Endonucleases were purchased from New England Biolabs (Ipswich, MA). ATRA was purchased from Sigma (St. Louis, MS) and used dissolved in DMSO. The transfection control plasmid pRL-TK (thymidine kinase promoter of Herpes Simplex virus with Renilla luciferase reporter gene) was purchased from Promega (Madison, WI). Mouse anti-human RARA and mouse anti-human RARB antibodies were purchased from Biomol International Inc. (Plymouth Meeting, PA). Rabbit anti-human RARG was purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA).

RNA isolation and reverse transcription

Total RNA from D407 cells was isolated with Qiagen RNeasy mini kit using a QiaCube instrument (Qiagen, Valencia, CA) following the manufacturer’s protocol. cDNA was synthesized from 2 μg of total RNA previously treated with DNase I (Invitrogen Corp, Carlsbad, CA) in a 20 μl reaction using SuperScript III Reverse Transcriptase (Invitrogen Corp, Carlsbad, CA).

Real time semi-quantitative RT-PCR

Real time semi-quantitative RT-PCR (qRT-PCR) was performed using SYBR green in an ABI 7500 instrument (Applied Biosystems Inc., Foster City, CA). A serial dilution of cDNA from the same source as
samples was used to obtain a calibration curve. Targets were quantified by determining the cycle threshold (Ct) and using the calibration curves. The relative values were normalized with 18S ribosomal RNA. Primers to quantify all msrAs were designed between exons 5 and 6 and primers for msrA2/3 transcripts were designed on exon 2 (Table 1, and reference 16).

Preparation of luciferase reporter constructs

Different areas of promoters 1 and 2 were amplified from human genomic DNA beginning at the transcription start site to different 5’ upstream region. The transcription start sites for all three transcripts were previously determined by 5’RACE [16]. The amplifications were performed using pfu DNA polymerase (Stratagene, La Jolla, CA). The PCR products were purified using the PCR purification Kit from Qiagen (Valencia, CA). The purified products were mixed with pGL4.10 reporter vector (Promega Corp, Madison, WI) which had been previously digested with EcoRV endonuclease and ligated using the Quick Ligase Kit (New England Biolabs, Ipswich, MA). Primers used to amplify promoters 1 and 2 are listed in the Table 1. Promoter sequences around 65 bp or less were cloned using adapter oligos with NheI and HindIII ends into pGL4.10 vector digested with the same endonucleases. All constructs were verified by direct sequencing.

Cloning of human RAR for expression

The open reading frame of RARA transcription factor was amplified by PCR from human retina cDNA using the forward primer, 5’-ACCATGGCCAGCAACAGCAG-3’ and the reverse primer, 5’-ATT-CACCGGGAGTGGGTGG-3’. PCR products were cloned into pcDNA 3.1/V5-His-TOPO vector (Invitrogen Corp, Carlsbad, CA) which drives expression using the CMV promoter.

DNA sequencing

Sequencing of the different plasmid constructs was performed using the BigDye terminator v3.1 cycle sequencing kit and an ABI 3130

Fig. 10. Effect of ATRA on the expression of native msrA transcripts. D407 cells were treated with 0, 1, 5, 10 and 15 μM of ATRA for 24 h and 48 h. Expression of msrA transcripts was determined by qRT-PCR using the same primers and conditions as Fig. 1 (Table 1). (A) All msrA transcripts (msrA1, 2 and 3) from both promoters. (B) MsrA transcripts from promoter 2 (msrA2/3). Cell viability (cellular dehydrogenase) was performed under the same treatment conditions. (C) Cell viability 24 h after treatment. (D) Cell viability 48 h after treatment. Values were normalized to the 18S ribosomal RNA and related to the control. Data are presented like mean±SD of at least three samples.
Genetic Analyzer instrument (Applied Biosystems Inc., Foster City, CA) following the manufacturer’s protocol.

Cell cultures

HEK 293 and SH-SY5Y cells were purchased from American Type Culture Collection (Manassas, VA). D407 cells were a kind gift from Dr. Richard Hunt (Department of Pharmacology and Microbiology, University of South Carolina, Columbia, SC). D407 were grown in DMEM medium supplemented with 4% fetal bovine serum (FBS). HEK293 were grown with DMEM supplemented with 10% FBS and SH-SY5Y neuroblastoma cells were grown with DMEM/F12 (1:1) supplemented with 10% FBS. Penicillin 10 U/mL, streptomycin 100 μg/mL and 2 mM of L-glutamine were added to all the cell media. DMEM and DMEM/F12 media were purchased from Atlanta Biologicals (Atlanta, GA) and all other components used for cell culture were from Invitrogen Corp (Carlsbad, CA).

Transient transfections and expression

The transfections were performed on 1×10^6 cells by electroporation using the Cell Line nucleofector V Kit and the Nucleofector™ II instrument (Axamta Biosystems Inc., Gaithersburg, MD) according to the manufacturer’s protocol. D407, SH-SY5Y and HEK293 cells were co-transfected with the different hMSRA-pGL4 constructs and a Renilla luciferase transfection control plasmid pRL-TK. Cells were plated on 24-well plates and fresh media was added 18 h after transfection. To determine the effects of RARA overexpression on the activity of MSRA promoters, D407 cells were transfected with 0.5 μg of the MSRA-pGL4 constructs, 1–2 or 2–3, 10 ng pRL-TK, and 4 μg of the expression plasmid RARA. The plasmid vector pCDNA 3.1/VS-His TOP0/IacZ (Invitrogen Corp, Carlsbad, CA) was used for the mock transfections.

Knockdown of RARA using small interference RNA (siRNA)

The RARA siRNA (cat# S100019369) and a negative control siRNA (cat# 1022083) were purchased from Qiagen Inc. (Valencia, CA). D407 cells (2×10^6 cells) were transfected with 500 nM of each siRNA as described above. The silencing of RARA siRNA was monitored by real time PCR using the forward primer, 5′-TGCCAGCTTACACATCCTCAT-3′ and the reverse primer, 5′-CTGCCACCCCTGCTGACCA-3′.

Cell viability assay

Cell viability was measured using the Cell Counting Kit-8 purchased from Dojindo Molecular Technologies, Inc. (Gaithersburg, MD). This assay measures cellular dehydrogenase activity. Absorbance was measured in a Wallac 1420 Victor 2 instrument (Perkin Elmer Inc, Waltham, MA).

Luciferase reporter assay

Luciferase expression was measured using the Dual-Luciferase Reporter Assay System (Promega) in a POLARStar OPTIMA Multifunction Microplate Reader instrument (BMG LABTECH Inc., Durham NC). The luciferase activity was quantified 48 h after transfection and normalized to the Renilla luciferase internal transfection control.

Treatment with all-trans retinoic acid

All experiments where ATRA was used were performed in dim yellow light to avoid photodestruction. In the studies where the effects of ATRA on MSRA promoter activity was measured (Figs. 6 and 8), ATRA (1, 5, 10 and 15 μM, final concentration) was added to the cells 18 h after transfection and the cells were harvested 24 h after ATRA treatment. In the studies where ATRA was used to induce msrA mRNA expression, 3×10^5 cells were seeded and the cells treated (in 6-well plates) for 24 and 48 h.

Immunobots

Nuclear extracts were prepared from D407 cells as previously described [23]. Nuclear proteins (15 μg per lane) were separated in 4–12% SDS-polyacrylamide gels (NuPAGE, Invitrogen Corp. Carlsbad, CA). Proteins were blotted onto nitrocellulose membranes using an iBlot dry blotting system (Invitrogen Corp.) following the manufacturer’s protocol. The membranes were blocked with 5% non-fat milk (BioRad Laboratories, Inc., Hercules, CA). The membranes were then incubated overnight with primary antibodies specific to RARA, RARB and RARG (see above). The blots were developed with anti-mouse (RARA and RARB) or anti-rabbit (RARG) horseradish conjugated secondary antibodies using Super Signal West Pico chemiluminescent substrate (Pierce Biotechnology Rockford, IL).

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Appendix A. Supplementary data

References


