

Depletion of Retinoic Acid Receptors Initiates a Novel Positive Feedback Mechanism that Promotes Teratogenic Increases in Retinoic Acid

Enrico D'Aniello¹, Ariel B. Rydeen^{1,2}, Jane L. Anderson¹, Amrita Mandal^{1,2}, Joshua S. Waxman^{1*}

1 The Heart Institute, Molecular Cardiovascular Biology and Developmental Biology Divisions, Cincinnati Children's Hospital Medical Center, Cincinnati, Ohio, United States of America, **2** Molecular and Developmental Biology Graduate Program, Cincinnati Children's Hospital Medical Center, Cincinnati, Ohio, United States of America

Abstract

Normal embryonic development and tissue homeostasis require precise levels of retinoic acid (RA) signaling. Despite the importance of appropriate embryonic RA signaling levels, the mechanisms underlying congenital defects due to perturbations of RA signaling are not completely understood. Here, we report that zebrafish embryos deficient for RA receptor α 1 (RAR α 1), a conserved RAR splice variant, have enlarged hearts with increased cardiomyocyte (CM) specification, which are surprisingly the consequence of increased RA signaling. Importantly, depletion of RAR α 2 or concurrent depletion of RAR α 1 and RAR α 2 also results in increased RA signaling, suggesting this effect is a broader consequence of RAR depletion. Concurrent depletion of RAR α 1 and Cyp26a1, an enzyme that facilitates degradation of RA, and employment of a novel transgenic RA sensor line support the hypothesis that the increases in RA signaling in RAR deficient embryos are the result of increased embryonic RA coupled with compensatory RAR expression. Our results support an intriguing novel mechanism by which depletion of RARs elicits a previously unrecognized positive feedback loop that can result in developmental defects due to teratogenic increases in embryonic RA.

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* E-mail: joshua.waxman@cchmc.org

Introduction

Improper retinoic acid (RA) signaling during development can cause congenital malformations that affect the forelimbs, ocular, cardiovascular, respiratory, urogenital and nervous systems [1–4]. Despite almost a century of investigation, the mechanisms underlying many congenital defects due to fluctuations in RA signaling are still not understood. RA acts as a ligand for RA receptors (RARs), members of the nuclear hormone family of transcription factors [5]. Work using disparate embryonic models has provided critical insight into the molecular mechanisms and developmental requirements of RAR function in vertebrate embryos [6–12]. In addition, RAR deficiency and inappropriate RA signaling are associated with numerous types of cancers [13]. In the majority of cases, the mechanism by which loss of RARs promote tumorigenesis is not understood. Therefore, understanding the roles of RARs during development will help elucidate the mechanisms underlying congenital defects, and possibly cancers, caused by inappropriate RA signaling [3,4].

RA signaling employs a number of feedback mechanisms in order to maintain appropriate levels in the embryo and tissues. The best characterized feedback mechanism is through regulation of the RA producing [retinol dehydrogenases (RDHs) and retinaldehyde dehydrogenases (Aldh1a)] and degrading (Cyp26) enzymes. Specifically, increased RA signaling inhibits the expression of the RA producing enzymes, while promot-

ing Cyp26a1 expression. Conversely, decreased RA signaling promotes expression of the RA producing enzymes, while inhibiting Cyp26a1 expression [14–18]. While modulation of RA signaling also affects the expression of other factors that control RA signaling [5,19], less well understood are feedback mechanisms that may influence RAR expression. RA response elements (RAREs) have been found in murine RAR α 2 and RAR β 2 promoters and RAR β 2 has been shown to be RA responsive [20–22]. However, if decreases in RA signaling, in particular due to loss of RAR expression, lead to compensatory expression of other RARs is less clear. While initial studies of mouse RAR KO mice suggested that there was not compensatory RAR expression in RAR deficient mice [11,12], more recent studies using siRNA to deplete RAR α have challenged this model and suggested that there may be compensatory RAR expression in RAR α deficient embryos [23]. Therefore, if there are RA feedback mechanisms that influence RAR expression and how the employment of these feedback mechanisms impact embryonic development are not well understood.

Here, we find that depletion of RAR α 1, a previously unrecognized yet conserved zebrafish RAR α splice variant, causes an increase in CM specification and heart size, which is due to the triggering of a feedback mechanism that surprisingly promotes increased RA signaling from surplus embryonic RA and compensatory RAR expression. Our results provide insight into a newly recognized positive feedback mechanism that we posit resists

Author Summary

Retinoic acid (RA) is the most active metabolic product of Vitamin A. Appropriate levels of RA are required for proper embryonic development and tissue maintenance in all vertebrates. Inappropriate levels of RA in human embryos can cause congenital defects that affect many organs, including the heart and limbs, and lead to numerous types of cancers. Understanding how animals maintain appropriate RA levels and the consequences of inappropriate RA signaling will therefore provide insight into human congenital defects and diseases. RA signaling is mediated by RA receptors (RARs), which are transcription factors that are activated when binding RA. We have found that depletion of RARs in zebrafish results in defects that are surprisingly due to increases in embryonic RA and not a deficiency of RA signaling. Our results are the first to demonstrate that RAR depletion elicits a positive feedback mechanism that promotes RA signaling through complementary increases in both embryonic RA and RAR expression. Therefore, our analysis provides novel insight into the molecular mechanisms that are required to maintain appropriate RA signaling and will positively impact our understanding of the mechanisms underlying congenital defects.

fluctuations in RA signaling due to perturbation in RAR expression. However, if improperly maintained, the positive feedback can result in RA induced congenital defects. Altogether, the results from this study significantly enhance our understanding of the feedback mechanisms that are used to maintain appropriate RA signaling levels and previously unexplored mechanisms that potentially underlie congenital defects.

Results

RAR α b1 deficient embryos have enlarged hearts and increased CM specification

In contrast to the studies of RARs in mice [9–12], depletion of RARs has not been able to recapitulate all of the consequences of loss of RA signaling in zebrafish [8], which prompted us to determine if additional conserved RAR variants exist in zebrafish beyond what has already been reported [24]. We cloned a previously unrecognized RAR α splice variant that is orthologous to human, mouse and *Xenopus* RAR α 1 termed RAR α b1 (Figure 1A–1C). The previously cloned zebrafish RAR α homologs RAR α a and RAR α b are teleost specific paralogs and both are orthologous to the splice variant 2 found in tetrapods (Figure 1B, 1D) [24]. Both *rar α b1* and *rar α b2* are expressed maternally and zygotically (Figure 1E), with ubiquitous expression until the tailbud stage (Figure S1A–S1I). After the tailbud stage, their expression patterns deviate (Figure 1F–1H and Figure S1J–S1O).

We used a translation blocking morpholino (MO) to examine the function of RAR α b1 (Figure 1B). By 48 hours post-fertilization (hpf), RAR α b1 deficient embryos had enlarged hearts with increased CM number and expression of CM marker genes *myl7*, *vmhc* and *amhc* (Figure 2A, 2B, 2M, 2N and Figure S2A–S2D). Similar increases in CM number were also found at 55 hpf (Figure S3A–S3C), suggesting the major addition of surplus CMs occurs during earlier stages of development. Consistent with this idea, we observed an expansion of CM differentiation (*myl7*, *vmhc*, and *amhc*) and progenitor (*nkx2.5* and *hand2*) marker expression in RAR α b1 deficient embryos at earlier stages in situ hybridization (ISH) and quantitative real-time PCR (qPCR; Figure 2C–2L,

2O–2Q). Injecting the RAR α b1 MO along with *rar α b1* mRNA that lacks the 5'UTR MO binding sequence is able to rescue the increased heart size, supporting the specificity of the phenotype (Figure S4A–S4D). Together, these results suggest that RAR α b1 deficient embryos have increased CM specification, number and heart size.

Depletion of zebrafish RAR α b paralogs promotes RA signaling

The increased atrial and ventricular CM number in RAR α b1 deficient embryos are reminiscent of RA signaling deficient embryos [25,26]. Therefore, we examined *hoxb5b* expression, which functions downstream of RA signaling to restrict atrial CM number [26] and is likely a direct target of RARs (Figure S5A–S5D). Unexpectedly, we found that *hoxb5b* expression was increased in RAR α b1 deficient embryos (Figure 3A–3C). While this was initially perplexing, our recent studies showed that *Hoxb5b* overexpression is able to mimic many of the teratogenic effects of RA treatment [27]. Therefore, we asked if the increases in *hoxb5b* expression in RAR α b1 deficient embryos could be a cause of the enlarged hearts. While depletion of *hoxb5b* alone using a low concentration of *hoxb5b* MO does not affect CM number (Figure S6A–S6C), we found that concurrent depletion of RAR α b1 and *Hoxb5b* largely restored heart morphology, CM differentiation marker expression, and CM number relative to the RAR α b1 deficient embryos (Figure 3F–3N), suggesting that the increased CM number in RAR α b1 deficient embryos is in part a consequence of the increased *hoxb5b* expression.

We next examined the expression of additional RA signaling responsive genes. Similar to *hoxb5b*, we found that the expression of additional RA signaling responsive genes, including *cyp26a1*, *dhrs3a*, *hoxb6b* and *hoxb5a*, was increased in RAR α b1 deficient embryos (Figure 3A). Comparing RA responsive gene expression in RA treated and RAR α b1 deficient embryos, we found that the trends were similar, but that RA treatment typically induced a greater increase in expression (Figure 3A). Conversely, treatment with DEAB, an antagonist of the RA producing enzyme Aldh1a, inhibited RA responsive gene expression (Figure 3A). These findings indicate that RAR α b1 depletion paradoxically results in increased expression of RA signaling responsive genes.

We next wanted to determine if increases in RA signaling responsive genes were specific to RAR α b1 depletion, so we examined RA responsive gene expression in RAR α b2 deficient embryos. Previous studies found that RAR α b2 deficient embryos lack forelimbs (pectoral fins) and *tbx5a* expression [8,28], which we confirmed (Figure S7A, S7C, S7D, S7F, S7H, S7I). However, similar to RAR α b1 depletion (Figure 3A and Figure 4A), RAR α b2 deficient embryos had increased expression of RA signaling responsive genes (Figure 4A). While the previous studies found a loss of forelimbs, defects in heart development were not reported. Despite the loss of forelimbs and increase in RA signaling responsive genes, we did not observe an increase in heart size, CM number or CM gene expression (Figure S8A–S8D). Therefore, although eliciting similar increases in RA signaling responsive gene expression, individual depletion of RAR α b1 and RAR α b2 results in distinct defects.

To determine the functional consequences of concurrent RAR α b1 and RAR α b2 depletion, we co-injected a suboptimal dose of each MO. Unfortunately, co-injection of an optimal dose of each MO resulted in significant non-specific toxicity even when injected along with *p53* MO. However, concurrent depletion of the RAR α s using suboptimal MO doses resulted in a dramatic increase in RA signaling responsive genes, above what was seen with depletion of RAR α b1 and RAR α b2 alone using the optimal

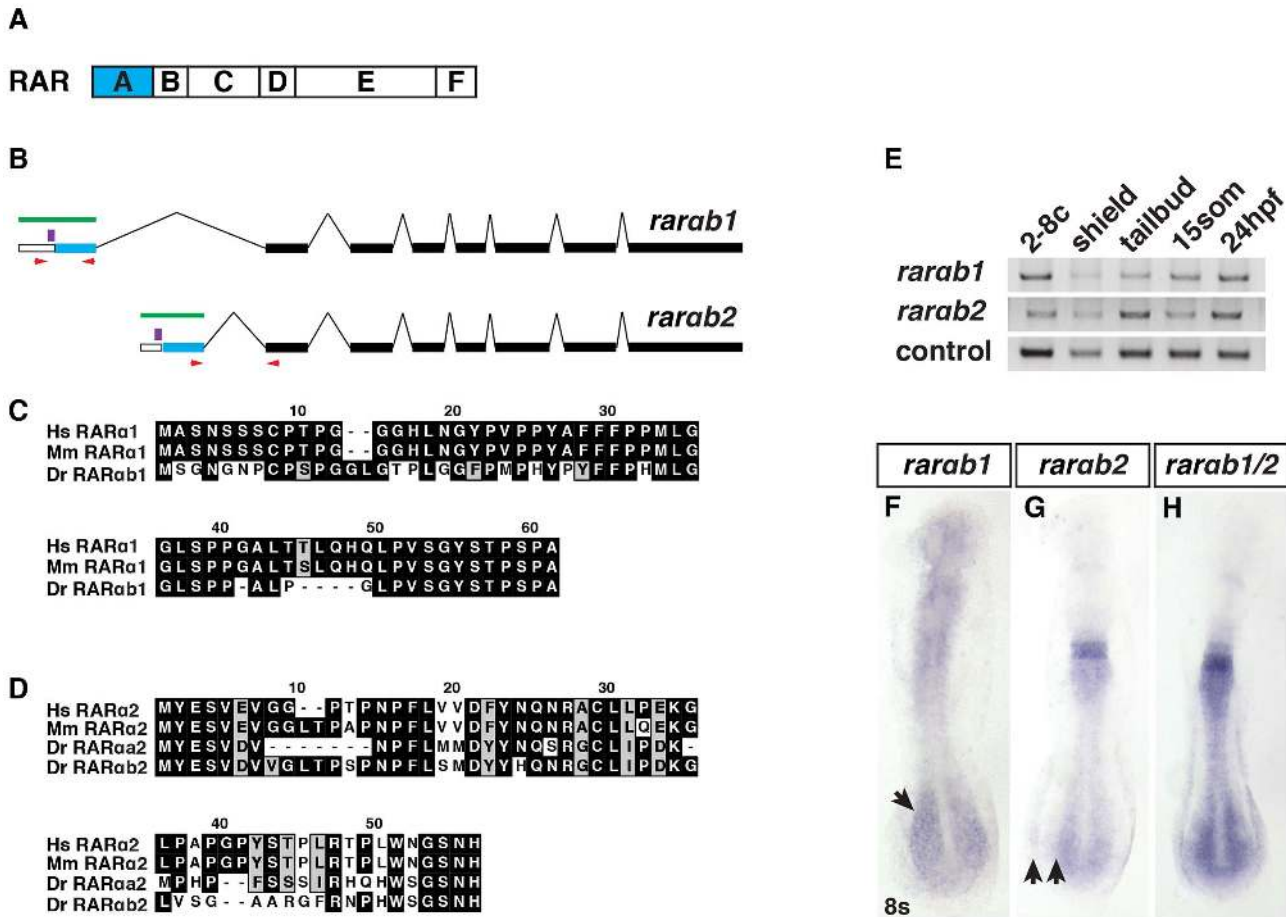


Figure 1. RARα1 and RARα2 sequences and expression. (A) Schematic representation of RAR domains. Blue box indicates the variable A domain, which is different between RARαb1 and the previously identified RARαb2 splice variant. (B) Schematic representation of RARαb1 and RARαb2 genomic organization (adapted from Ensemble_v9). Blue bars represent the first exon, which encodes the respective A domains. White bars represent the 5' UTRs. Black bars represent the exons that are common to the two variants. Green bars represent the target of the antisense probes used for ISH. Red arrows indicate the position of the primers used to perform RT-PCR. Purple bars indicate the position of the morpholino target sequences. (C) Alignments of the A domains of human (Hs) RARα1, mouse (Mm) RARα1, and zebrafish (Dr) RARα1. The presence of this previously unrecognized splice variant was recently confirmed in the latest zebrafish genome assembly (Ensemble Zv9). There is no RARαa splice variant 1 ortholog in the zebrafish genome. (D) Alignments of the A domains of Hs RARα2, Mm RARα2, Dr RARα2, and Dr RARαb2. (E) Reverse transcriptase PCR (RT-PCR) for the zebrafish *rarαb* isoforms. *max* was used as the control. -RT control did not reveal genomic contamination (data not shown). (F) *Rarαb1* is expressed in the ventral anterior of the embryo and the presomitic paraxial mesoderm (arrow) at the 8 somite (s) stage. (G) *Rarαb2* is expressed in rhombomeres 5 and 6, the spinal cord and the posterior lateral plate mesoderm (LPM). Arrows indicate the space between the posterior spinal cord and LPM expression domains. (H) Together, the expression patterns recapitulate a previously reported *rarαb* probe (referred to as *rarαb1/2*), which detects both isoforms [24]. In F–H, embryos are flatmounted and are dorsal views with anterior up. doi:10.1371/journal.pgen.1003689.g001

MO doses (Figure 4A). Additionally, there was an anterior shift of *hoxb5a* expression in the spinal cord of RARαb1+2 deficient embryos, suggesting the spinal cords are posteriorized (Figure S9A–S9E). Increased RA signaling inhibits *aldh1a2* expression through a negative feedback mechanism [16–18]. Although *aldh1a2* expression in individual RARαb1 and RARαb2 deficient embryos was not suppressed (Figure 4B), *aldh1a2* expression was decreased in embryos depleted for both RARαb variants (Figure 4B). To corroborate the increases in endogenous RA signaling responsive genes, we used the RA signaling reporter line *Tg(12XRARE-ef1a:EGFP)^{sk72}* [29]. Again, co-depletion of both RARαbs resulted in a greater expansion of *egfp* expression, compared to the individual depletion of each RARαb (Figure 4D–4H). Therefore, these experiments support the hypothesis that the RARαb1+2 deficient embryos are sensing more significant increases in RA signaling than embryos deficient for either RARαb variant alone.

We next examined the consequences of this functional interaction on heart development. We found that the hearts of RARαb1+2 deficient embryos had increased atrial size, CM number, and a dramatic increase in *amhc* expression (Figure 4I, 4L–4N and Figure S10A–S10D). Significant effects on CM number or heart size were not found when using a suboptimal dose of either RARαb1 or RARαb2 MO alone (Figure 4I–4K, 4M), though we did find a modest increase in CM marker gene expression in the RARαb1 deficient embryos (Figure 4N). Interestingly, in RARαb1+2 deficient embryos we found more significant increases in atrial CM number and *amhc* expression (Figure 4M, 4N), which were remarkably similar to the consequences of modest increases in RA signaling due to RA treatment [27]. Increased RA signaling can also inhibit forelimb development [17] and RARαb1 deficient embryos also have smaller forelimbs and a modest reduction of *tbx5a* expression (Figure S7A, S7B, S7D, S7F, S7G, S7I). A functional interaction

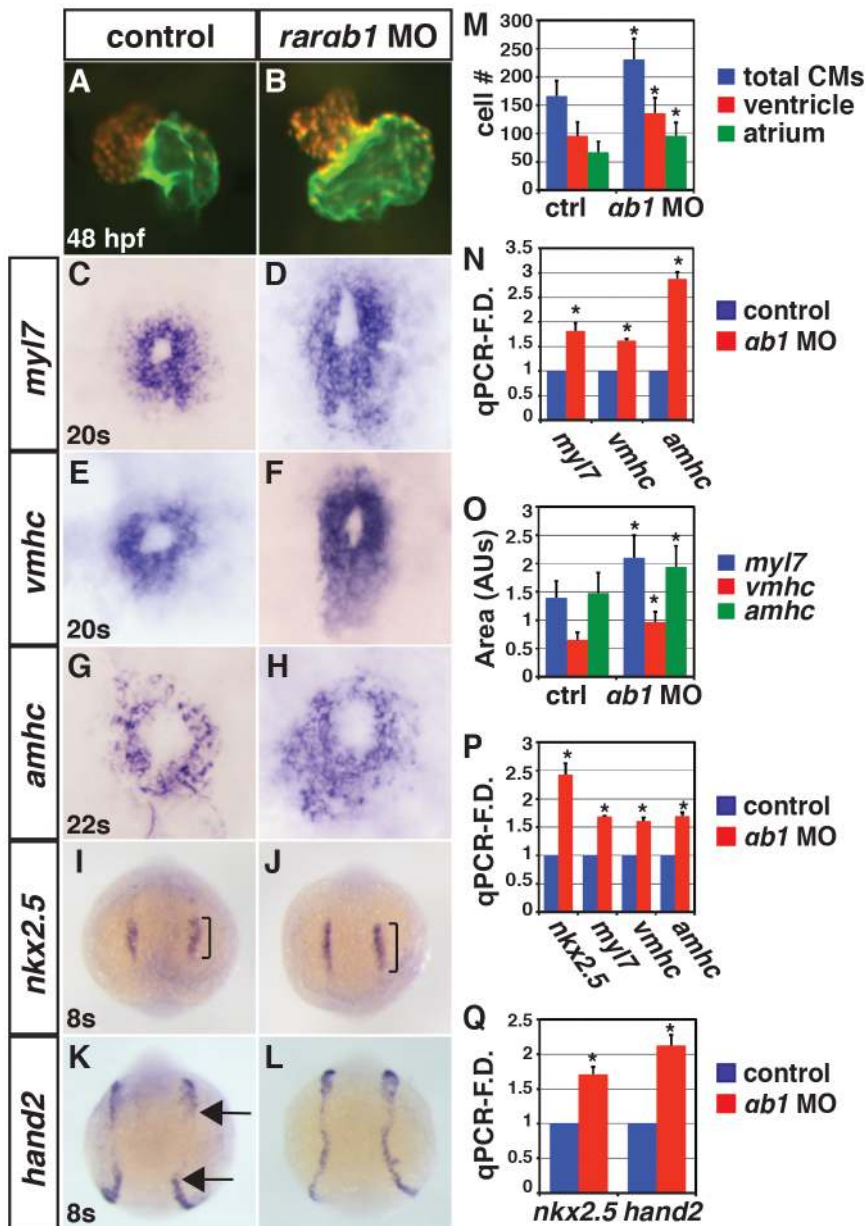


Figure 2. RAR α 1 deficient embryos have enlarged hearts with increased CM number. (A, B) Hearts from control sibling and RAR α 1 deficient *Tg(-5.1myl7;DsRed-NLS)²* embryos. Images are frontal views. Red indicates ventricle. Green indicates atrium. (C–H) ISH for CM differentiation marker genes. (I–L) ISH for CM progenitor marker genes. Brackets in I and J indicate length of *nkx2.5* expression. Arrows in K indicate posterior and anterior limits of the *hand2* expression domains in the LPM. In C–L, views are dorsal with anterior up. (M) Mean CM number at 48 hpf. (N) qPCR for CM differentiation marker gene expression at 48 hpf. (O) Areas of the amount of cells expressing the CM differentiation marker genes at the 20 s and 22 s stages. (P) qPCR for CM differentiation marker gene and *nkx2.5* expression at 24 hpf. (Q) qPCR for CM progenitor gene expression at the 8 s stage. Asterisk in all graphs indicate a statistically significant difference compared to controls ($p < 0.05$). Error bars in all graphs indicate standard deviation. doi:10.1371/journal.pgen.1003689.g002

with the RAR α variants that resulted in loss of forelimbs was also observed (Figure S7D, S7E). Therefore, concurrent depletion of RAR α variants elicits increases in RA signaling with heart and forelimb phenotypes that are strikingly similar to increases in RA signaling caused from RA treatment.

RAR α 1 deficient embryos have increased embryonic RA

We sought to understand the mechanism underlying the increase in RA signaling in RAR α deficient embryos. In the absence of RA, RARs are thought to interact with transcriptional

co-repressors, while binding of RA converts the RARs to transcriptional activators [1,5]. A previous study in *Xenopus* suggested that RARs are required as transcriptional repressors in some developmental contexts [6]. However, our gain-of-function analysis did not support that these zebrafish RARs function as transcriptional repressors (Figure S11A–S11L), consistent with what we have reported previously [29]. However, Manshour et al. [23] found a compensatory increase in the expression of other RARs when using siRNA to deplete RAR α in mice. Similarly, we found that the expression of other zebrafish RARs [24] was

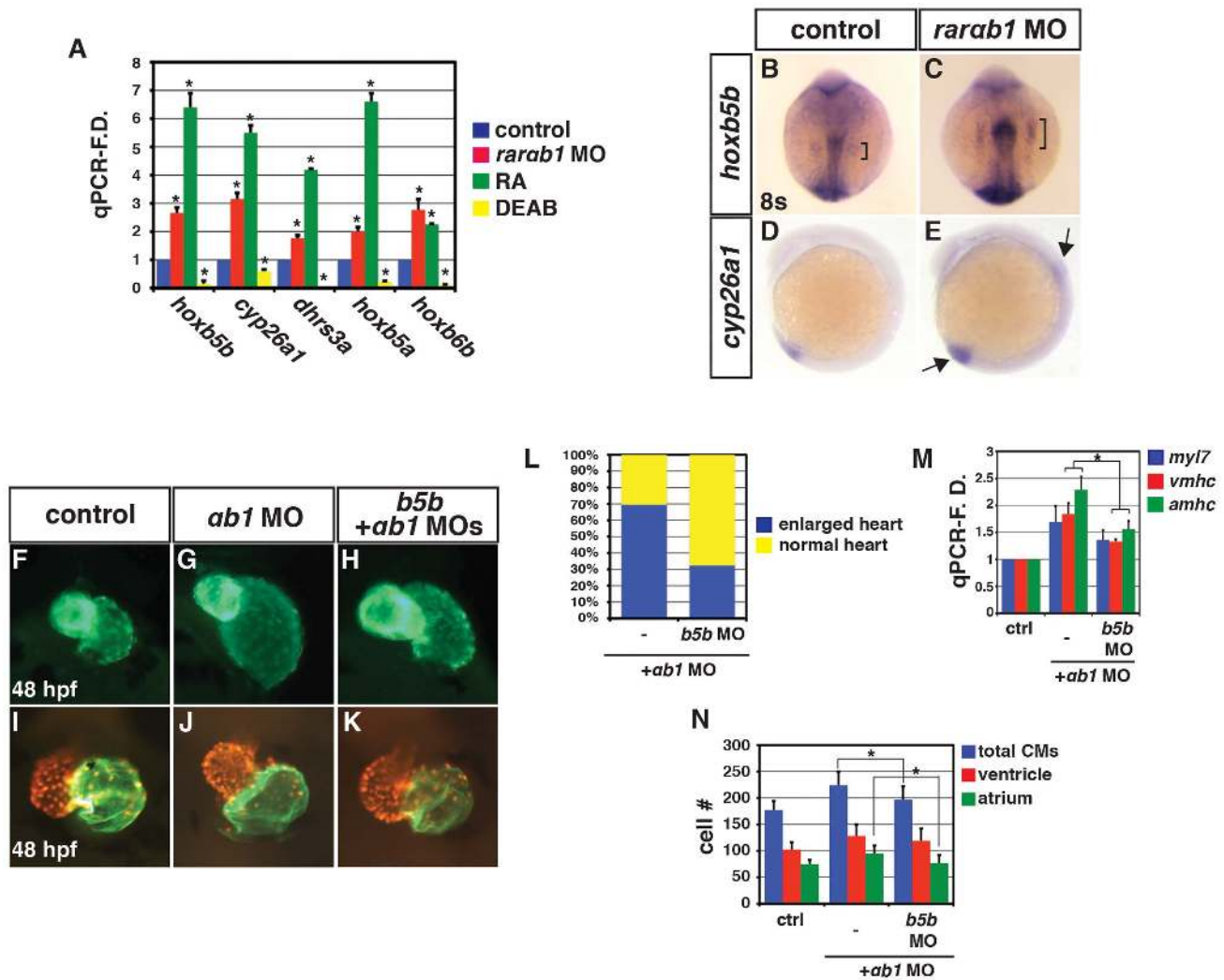


Figure 3. RAR α b1 deficient embryos have increased expression of RA signaling responsive genes. (A) qPCR for RA signaling responsive gene expression at the 8 s stage. (B, C) ISH for *hoxb5b* expression at the 8 s stage. Bracket indicates length of expression in the LPM. Views are dorsal with anterior up. (D, E) ISH for *cyp26a1* expression at the 8 s stage. Arrows in E indicate increased expression in the tailbud and spinal cord. Views are lateral with anterior up and dorsal right. (F–H) Fronto-lateral views of *Tg(-5.1myl7:GFP)²* embryos at 48 hpf of control sibling, RAR α b1 deficient embryos, and RAR α b1+Hoxb5b deficient embryos. (I–K) Hearts from control sibling, RAR α b1 deficient embryos and RAR α b1+Hoxb5b deficient *Tg(-5.1myl7:DsRed-NLS)²* embryos. Images are frontal views. Red indicates ventricle. Green indicates atrium. (L) Percentage of control+*rarab1* MOs (n = 60), *hoxb5b*+*rarab1* MOs (n = 68) showing enlarged and normal hearts. (M) qPCR for CM differentiation gene expression at 48 hpf. (N) Mean CM number at 48 hpf.

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increased in RAR α b deficient embryos (Figure 4C and Figure S12A–S12L), suggesting that compensatory RAR expression is a conserved response to depletion of RAR α homologs in vertebrates. Although Manshoury et al. [23] proposed the compensatory RAR expression was RA signaling dependent, our results suggest that the expression of most RARs is potentially regulated independent of RA signaling (Figure 4C), because the effects on RAR expression did not parallel modulation of RA signaling using RA and DEAB. While we observed compensatory expression of other RARs in RAR α b deficient embryos, it is difficult to conclude that increased RAR expression is the sole cause of the increase in RA signaling since overexpression of RARs in zebrafish embryos does not produce significant positive or negative effects on RA responsive gene expression (Figure S11A–S11J) [29]. Nevertheless, our results suggest that when depleting RAR α bs in zebrafish embryos compensatory RARs are present that can mediate RA signaling.

Because we did not have evidence that RARs act as transcriptional repressors or that the increased expression of RARs alone contributes to the increases in RA signaling in RAR α b deficient embryos, we hypothesized that the depletion of RARs may trigger an increase in embryonic RA. Although *aldh1a2* expression was suppressed in RAR α b1+2 deficient embryos similar to when embryos sense increases in RA signaling (Figure 4B) [16–18], the expression of *rdh10a* and *rdh10b*, which control a limiting step in RA production in vertebrates by generating retinal from retinol [14,15], was increased in RAR α b1 and RAR α b1+2 depleted embryos (Figure 4B and Fig. S13A–S13C). Interestingly, *rdh10b* expression, which was not sensitive to modulation of RA signaling, was increased in RAR α b deficient embryos (Figure 4B). Therefore, our results suggest that depletion of RAR α bs triggers an increase in RA through promoting *rdh10* expression.

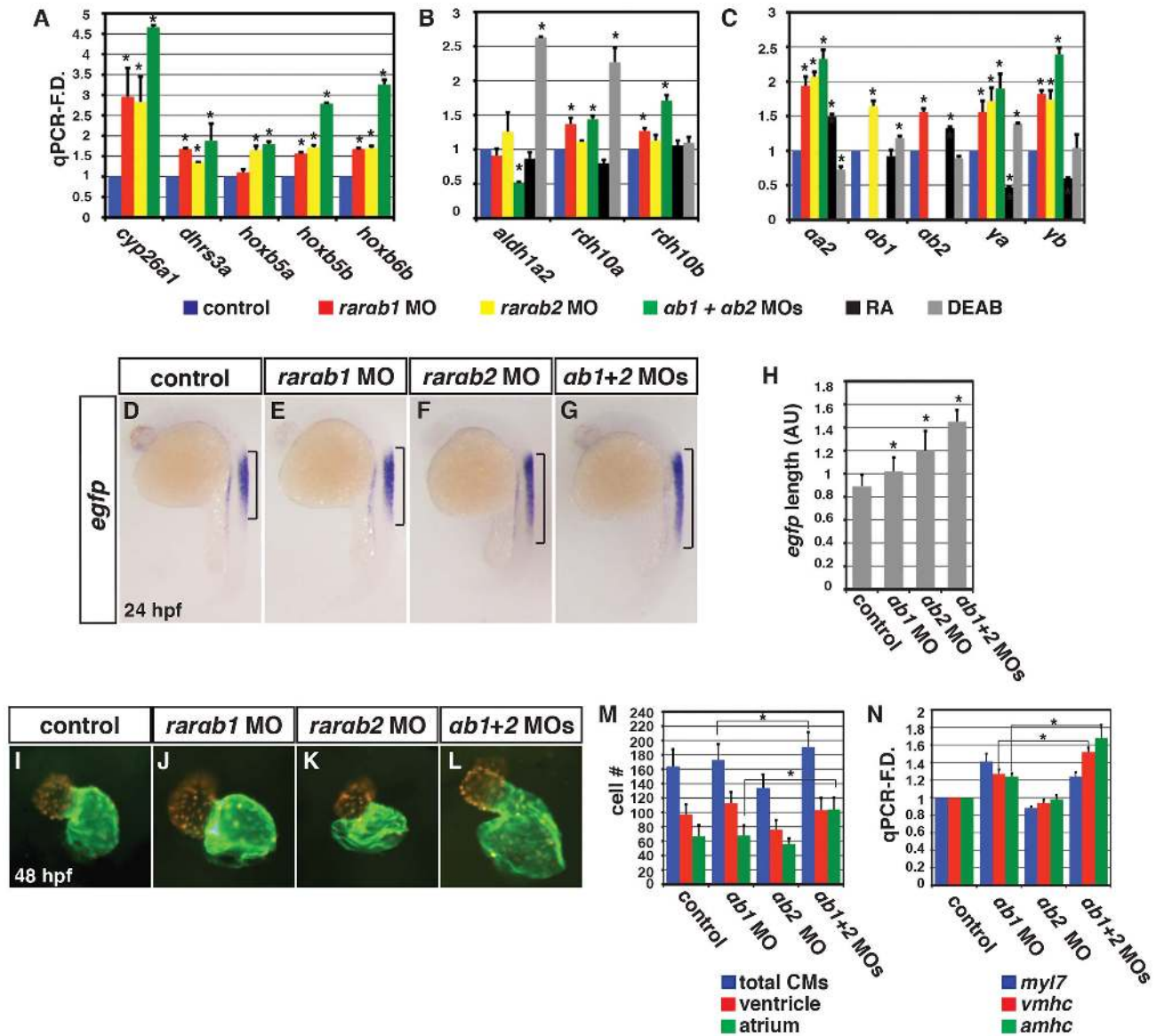


Figure 4. Concurrent depletion of RAR α 1 and RAR α 2 promotes increased RA signaling and atrial CM number. qPCR for (A) RA signaling responsive gene, (B) RA metabolizing gene, and (C) zebrafish *rar* expression in control sibling, RAR α 1 deficient, RAR α 2 deficient, RAR α 1+RAR α 2 (suboptimal doses) deficient, RA treated, and DEAB treated embryos at the 8 s stage. (D–G) ISH for *egfp* expression in *Tg(12XRARE-ef1a:EGFP)^{sk72}* embryos. Brackets indicate the length of *egfp* expression in the spinal cord. (H) Measurements of the length in arbitrary units (AU) of *egfp* expression in the spinal cord of *Tg(12XRARE-ef1a:EGFP)^{sk72}* embryos. (I–L) Hearts from control and RAR α 1 and RAR α 2 deficient *Tg(-5.1myl7:DsRed-NLS)^{f2}* embryos. Images are frontal views. Red indicates ventricle. Green indicates atrium. (M) Mean CM number from *Tg(-5.1myl7:DsRed-NLS)^{f2}* hearts at 48 hpf. (N) qPCR for CM marker gene expression at 48 hpf. While modest increases in *vmhc* expression in RAR α 1+RAR α 2 deficient embryos were observed relative to RAR α 1 (suboptimal dose) deficient embryos, corresponding increases in ventricular CM number were not observed. doi:10.1371/journal.pgen.1003689.g004

In addition to inhibiting *aldh1a2* expression, increased RA signaling promotes a negative feedback mechanism that limits RA levels by positively regulating *Cyp26a1* expression [16–18]. Since we observe an increase in *cyp26a1* expression in RAR α 1 deficient embryos (Figure 3A, 3D, 3E and Figure 4A), which was also consistent with the hypothesis that there is increased embryonic RA, we postulated that the increased *Cyp26a1* may be protecting the RAR α 1 deficient embryos from teratogenic increases in embryonic RA. Therefore, we concurrently depleted RAR α 1 and *Cyp26a1* to determine if there was a functional interaction indicative of increased embryonic RA. For these experiments, a suboptimal dose of *cyp26a1* MOs (Figure S14A–

S14E) was used to more easily discern a functional interaction. In either the RAR α 1 or *Cyp26a1* deficient embryos alone, we never observed absence of the MHB or defects in tail elongation (Figure 5A–5C, 5E–5G). However, co-depletion of RAR α 1 and *Cyp26a1* resulted in a loss of the MHB and truncated tails (Figure 5D, 5H), similar to increases in RA signaling [17,19,29,30]. Furthermore, we found that RAR α 1+*Cyp26a1* deficient embryos had dismorphic hearts with a specific reduction in ventricular CM number compared to controls embryos hearts (Figure 5I–5L, 5Q), which interestingly resembles the trend we previously found in embryos with intermediate increases in RA signaling [27].

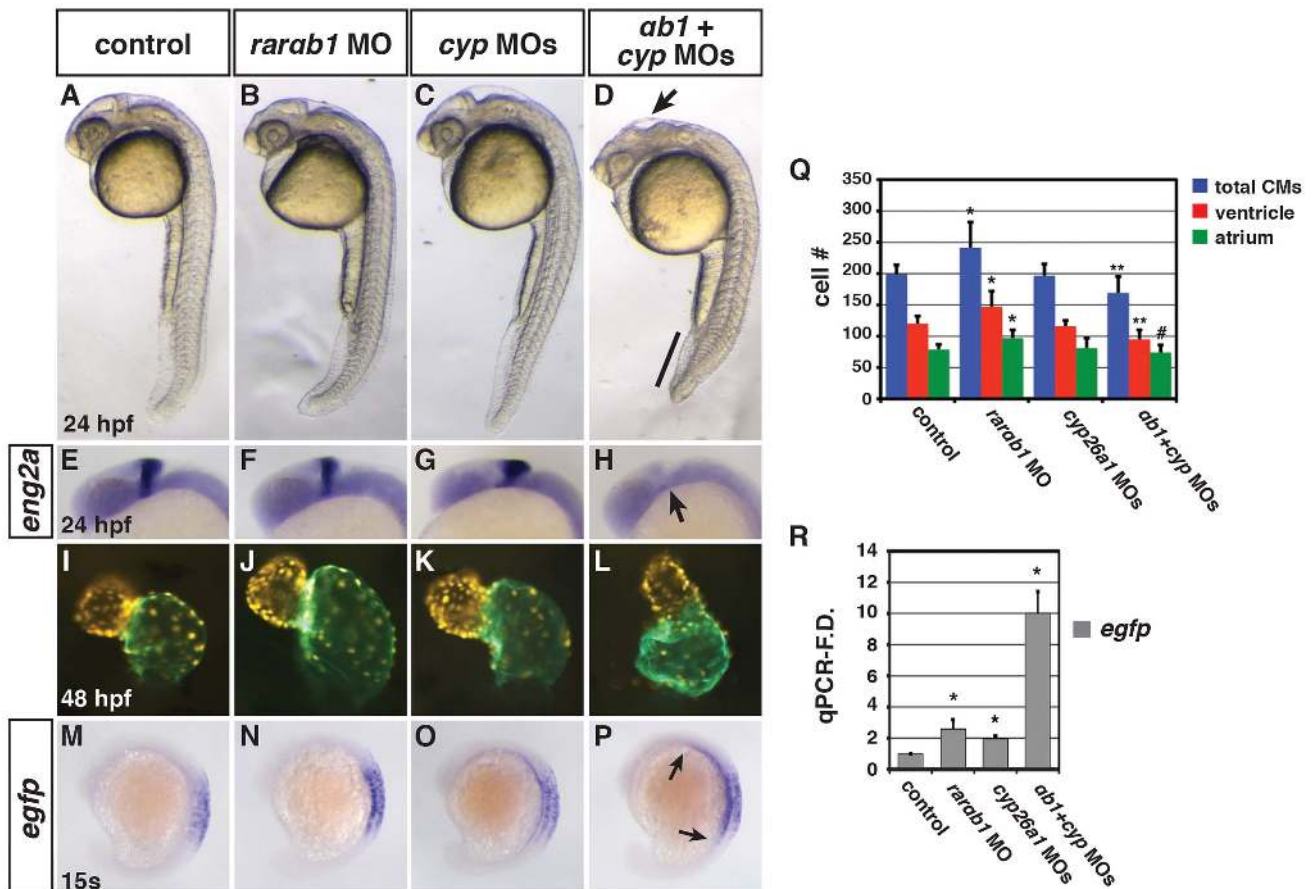


Figure 5. Concurrent depletion of RARαb1 and Cyp26a1 results in phenotypes resembling RA treatment. (A–D) Control sibling, RARαb1 deficient, Cyp26a1 deficient, and RARαb1+Cyp26a1 deficient embryos. A suboptimal dose of the *cyp26a1* MOs was used that did not cause ostensible defects for these experiments. In D, arrow indicates loss of the MHB and line indicates shortened tail. Images are lateral views with dorsal right and anterior up. (E–H) ISH for *eng2a*, which marks the MHB. 100% of (E) control sibling (n=11), (F) RARαb1 deficient (n=7), and (G) Cyp26a1 deficient (n=7) had *eng2a* expression. 85% of (H) RARαb1+Cyp26a1 deficient embryos (n=7) had a complete absence of *eng2a* expression (arrow in H). Equivalent results were obtained using *pax2a*, which also marks the MHB (data not shown). (I–L) Hearts from control sibling, RARαb1 deficient, Cyp26a1 deficient, and RARαb1+Cyp26a1 deficient *Tg(-5.1myl7:DsRed-NLS)^{f2}* embryos. Images are lateral views. Red indicates ventricle. Green indicates atrium. (M–P) ISH for *egfp* in *Tg(β-actin:GDBD-RLBD)^{ccH1}; Tg(UAS:EGFP)* embryos. Lateral views with dorsal right and anterior up. (Q) Mean CM number at 48 hpf and (R) qPCR for *egfp* expression at 15 s in control sibling, RARαb1 deficient, Cyp26a1 deficient, and RARαb1+Cyp26a1 deficient embryos. Double asterisks in Q indicate a statistically significant difference relative to control and RARαb1 deficient embryos. Pound sign in Q indicates a statistically significant difference relative to RARαb1 deficient embryos.

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Although one interpretation of the functional interaction of RARαb1 and Cyp26a1 depletion is that there is increased embryonic RA levels in these embryos, we wanted to further test this hypothesis using additional assays. First, we sought to use a distinct readout of embryonic RA, so we made a novel stable transgenic RA sensor line which incorporated the RARαb ligand binding domain (RLBD) fused to the Gal4 DNA binding domain (GDBD) expressed under the β-actin promoter (Figure S15A–S15G) [31]. Previous studies have found that similar GDBD fusions with nuclear hormone receptor LBDs create an effective reporter of nuclear hormone activity [6,32,33]. We observed a dramatic increase in reporter expression when RARαb1 and Cyp26a1 were depleted together in *Tg(β-actin:GDBD-RLBD); Tg(UAS:EGFP)* embryos (Figure 5M–5P, 5R) [34]. Second, our hypothesis predicted that reducing embryonic RA levels should be able to rescue teratogenic phenotypes found in RARαb1+Cyp26a1 and RARαb1 deficient embryos. Consistent with this hypothesis, DEAB treatment of RARαb1+Cyp26a1 deficient embryos was able to rescue the loss of MHB (Figure 6A–

6J). Additionally, treatment of RARαb1 deficient embryos with DEAB partially rescue the enlarged heart phenotype and restored atrial CM number (Figure 6K–6O). Lastly, our hypothesis predicts that exogenous treatment with a concentration of RA that causes a minor increase in RA signaling should result in aberrant heart phenotypes that are similar to RARαb1 deficient embryos. Indeed, embryos treated with low concentrations of exogenous RA (lower than we had reported using previously [27]) had enlarged hearts with an increase in both atrial and ventricular CM number at 48 hpf (Figure 6P–6R). Altogether, our results suggest that increases in embryonic RA, coupled with compensatory RAR expression, contribute to the developmental defects found in RARαb1 deficient embryos.

Discussion

Together, our study supports a novel paradigm whereby RARαb depletion elicits a positive feedback mechanism that can result in teratogenic increases in RA signaling. Importantly, our

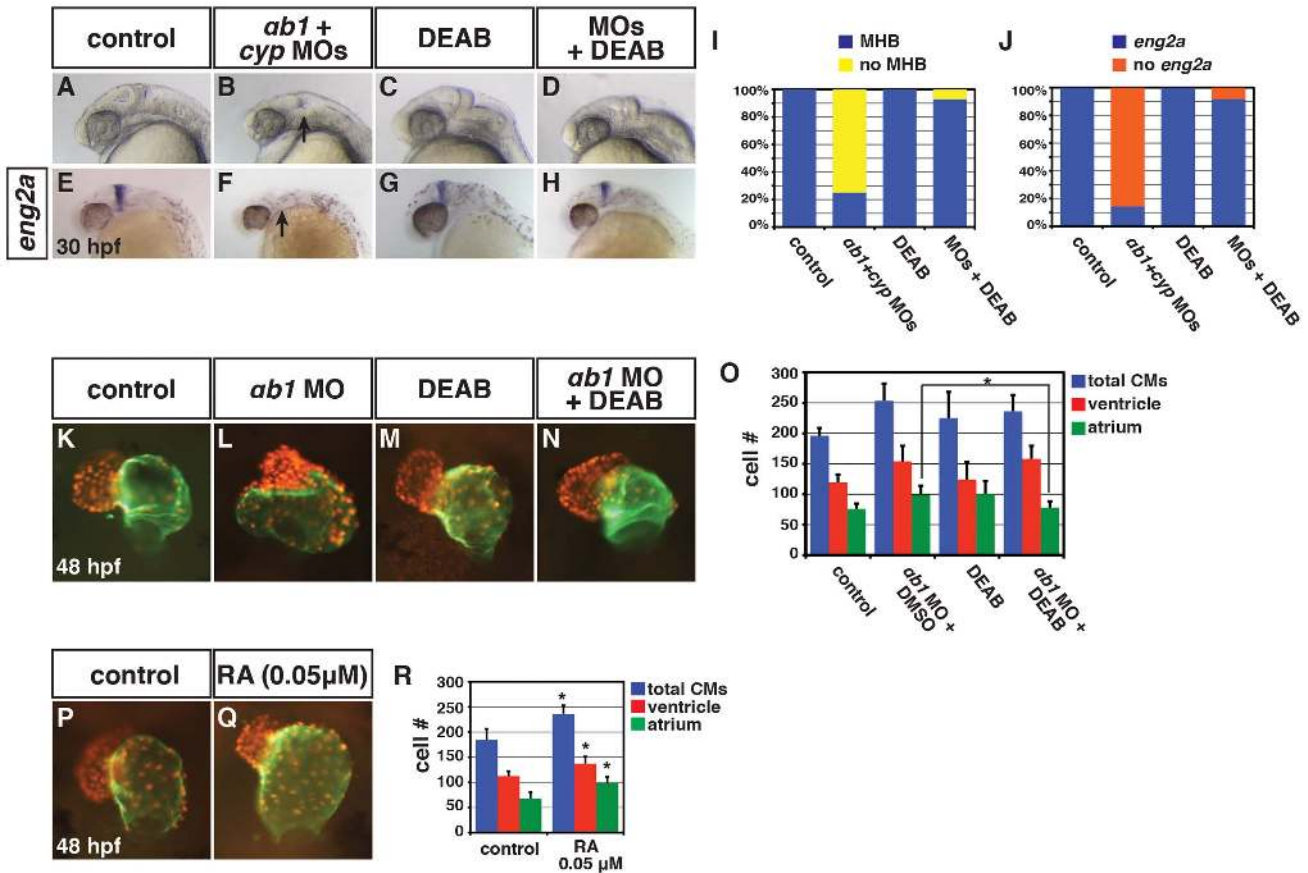


Figure 6. Reduction of RA in RAR α b1 deficient embryos can rescue developmental defects. (A–H) Control sibling, RAR α b1+Cyp26a1 deficient, control sibling treated with DEAB, and RAR α b1+Cyp26a1 treated with DEAB embryos. In B and F, arrows indicates loss of the MHB and *eng2a* expression. Images are lateral views with dorsal right and anterior up. (I) Percentage of control sibling (n = 16), RAR α b1+Cyp26a1 deficient embryos (n = 16), control sibling embryos treated with DEAB (n = 13), and RAR α b1+Cyp26a1 deficient embryos treated with DEAB (n = 14) that had a MHB based on morphology. (J) Percentage of control sibling (n = 17), RAR α b1+Cyp26a1 deficient embryos treated with DEAB (n = 15), and RAR α b1+Cyp26a1 deficient embryos treated with DEAB (n = 12) that had *eng2a* expression at the MHB. (K–N) Hearts from *Tg(-5.1myl7:DsRed-NLS)^{f2}* control sibling, RAR α b1 deficient, DEAB treated, and DEAB+RAR α b1 deficient embryos. Images are frontal views. Red indicates ventricle. Green indicates atrium. (O) Mean CM number at 48 hpf. (P,Q) Hearts from *Tg(-5.1myl7:DsRed-NLS)^{f2}* control sibling embryos and *Tg(-5.1myl7:DsRed-NLS)^{f2}* embryos treated with a low concentration of RA. Images are frontal views. Red indicates ventricle. Green indicates atrium. (R) Mean CM number at 48 hpf. doi:10.1371/journal.pgen.1003689.g006

work highlights that loss and gain of RA signaling can cause similar developmental defects. RA signaling is required to restrict CM specification [25,26], while high increases in RA signaling can eliminate CM specification (Figure 7A) [27]. However, our present findings suggest that low increases in RA signaling, achieved when treating embryos with μ M concentrations of RA or through RAR α b depletion, can also promote increases in both atrial and ventricular CM specification (Figure 7A). As we found previously, modest, but slightly higher increases of RA signaling can promote atrial CM specification without significantly affecting ventricular CM specification [27], which is strikingly similar to what we found with concurrent depletion of the RAR α b variants here (Figure 7A). Moreover, intermediate increases in RA signaling can inhibit ventricular CM specification, which is similar what we observed when concurrently depleting RAR α b1 and Cyp26a1 (Figure 7A). It also appears that modulation of Hox activity downstream of both gain and loss RA signaling is at least partially responsible for the increases in CM specification, suggesting the hypothesis that the similar effects on CM number are actually due to opposite perturbations of anterior-posterior patterning within the ALPM. Therefore, our analysis corroborates and extends previous

observations that there are differential effects on atrial and ventricular CM populations as there is a progressive increase from low to intermediate levels of RA signaling in the early embryo.

It is interesting that depletion of RAR α homologs using MOs in zebrafish, presented in this study, and *Xenopus* [6] elicit similar phenotypic responses. In *Xenopus* embryos, RAR α depletion alone results in loss of the MHB [6]. While depletion of RAR α b1 alone does not result in MHB defects in zebrafish embryos, we have found that RAR α b1+Cyp26a1 deficient embryos completely lack the MHB. Taken together, these results suggest that the underlying consequences of increased RA signaling due to depletion of RAR α homologs are likely conserved at least in *Xenopus* and zebrafish embryos, but that in *Xenopus* perhaps the role of Cyp26 enzymes in protecting the brain has been lost. Despite similarities in the phenotypes that both point to an increase in RA signaling in RAR α and RAR α b deficient *Xenopus* and zebrafish embryos, our results contrast with the model proposed by Koide et al. [6], which concluded that RARs are required to function as transcriptional repressors. Importantly, the tools used in the previous study, including dominant-negative RARs, transcriptional co-repressors, and

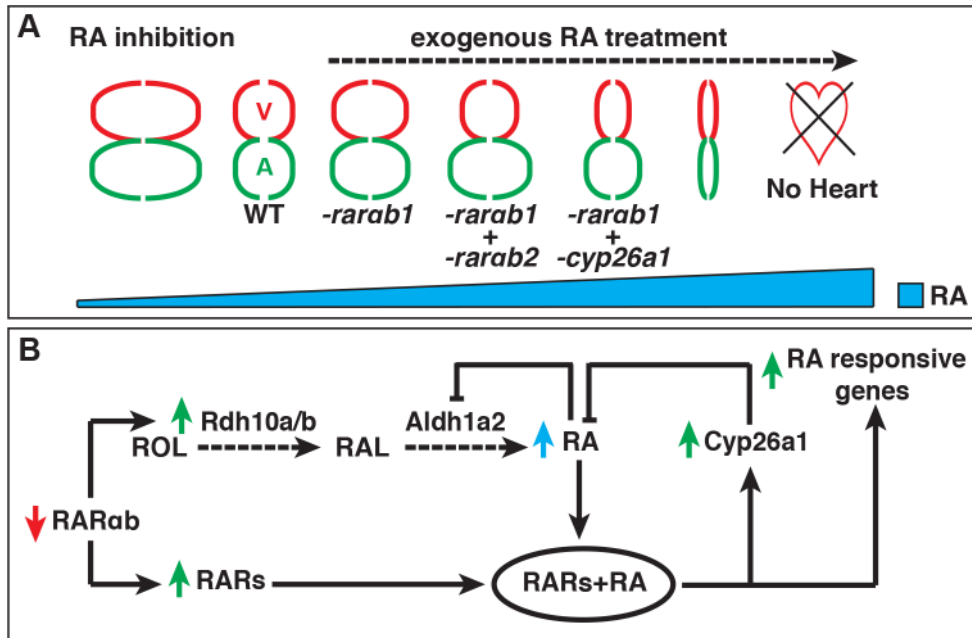


Figure 7. Models of the effects of RA signaling on heart patterning and the RA feedback mechanism. (A) Model depicting the consequences on atrial and ventricular CM specification at different levels of RA signaling. (B) Model of the previously unrecognized feedback mechanism that triggers increased RA signaling when depleting RARs. ROL = retinol. RAL = retinal. Red and green arrows indicate the effects on gene expression. Blue arrow indicates the effect on RA levels. doi:10.1371/journal.pgen.1003689.g007

inverse agonists, would not have allowed them to distinguish between a transcriptional de-repressive model and the positive feedback mechanism involving the production of excess RA supported here.

In addition to the phenotypic similarities when depleting RAR α homologs in *Xenopus* and zebrafish, depletion of zebrafish RAR α s results in compensatory RAR expression similar to RAR α depletion in mice [23], supporting the hypothesis that this feedback response to RAR α deficiency is conserved in vertebrates. Importantly, the response to RAR depletion is likely different than complete ablation of RARs. RAR KO mice have not been reported to have compensatory increases in other RARs [11,12], suggesting that a complete loss of RAR expression may cause a breakdown of this feedback loop. However, when considering the probability that RAR expression would be completely lost vs. depleted, we postulate that insults resulting in depletion of RAR expression would be much more likely. Consistent with this idea, variable levels of RAR expression deficiency, which in the case of RAR β can be due to epigenetic silencing, is commonly observed in a variety of cancers [13].

Given the conserved feedback mechanisms already recognized that limit fluctuations in RA signaling in vertebrates [16,17,19,23], it seems logical that a conserved mechanism that senses RAR deficiency would also exist to prevent loss of RA signaling. We propose that this newly recognized positive feedback mechanism would be more suitable to prevent transient deficiency in RARs. As demonstrated here, persistent RAR α b depletion can result in a hypervigilant response of RA signaling and RA-induced teratogenic defects. Overall, these data provide insight into a previously unappreciated RAR-dependent positive feedback mechanism (Figure 7B), which is active during development. Further elucidation of this RA signaling feedback mechanism may illuminate the etiology of poorly understood RA-insensitive cancers [13,23] and congenital defects [1,3].

Materials and Methods

Ethics Statement

All zebrafish husbandry and experiments were performed in accordance with protocols approved by the Institutional Animal Care and Use Committee (IACUC) of Cincinnati Children's Hospital Medical Center.

Zebrafish husbandry and transgenic lines

Zebrafish (*Danio rerio*) were raised and maintained as previously described [35]. The following transgenic lines were used: *Tg(-5.1myl7:DsRed-NLS)* [36], *Tg(-5.1myl7:EGFP)^{tcu26}* [37], *Tg(12XRARE-ef1a:EGFP)^{sk72}* [29], *Tg(β -actin:GDBD-RLBD)^{ech1}* (was created using the Gateway/Tol2 system [38] and additional characterization is reported in [31]), *Tg(UAS:EGFP)* [34], and *Tg(UAS:nfsB-mcherry)* [39].

ISH

Whole-mount ISH was carried out using standard procedures [40]. All probes except *rarb1* (accession number: KF030797) and *rarb2* were reported previously. *myl7* (formerly called *cmlc2*; ZDB-GENE-991019-3), *amhc* (ZDB-GENE-031112-1), *vmhc* (ZDB-GENE-991123-5), *nkx2.5* (ZDB-GENE-980526-321), *hand2* (ZDB-GENE-000511-1), *hoxb5a* (ZDB-GENE-980526-70), *hoxb5b* (ZDB-GENE-000823-6), *dhx3a* (ZDB-GENE-040801-217), *cyp26a1* (ZDB-GENE-990415-44), *rarb1/2* (which recognizes both isoforms and was formerly called *rarb* [24]; ZDB-GENE-980526-72), *rara* (ZDB-GENE-980526-284), *rarya* (ZDB-GENE-980526-531), *raryb* (ZDB-GENE-070314-1), *rdh10a* (ZDB-GENE-070112-2242), *tbx5a* (ZDB-GENE-030909-7), *eng2a* (ZDB-GENE-980526-167), *egr2b* (formerly called *krox20*; ZDB-GENE-980526-283), *egfp* (accession number: JQ064510.1), and *mcherry* (accession number: JN795134.1).

MO and mRNA injections

The *rarb1* MO (5'-TGCAGGTCATCCGTAATGCCGATC) was designed to the 5' UTR of *rarb1*. Additional MOs targeting another region of the 5' UTR and the donor splice junction, which saturated the available MO target sites, were also tried. However, injection of these MOs resulted in significant toxicity and were not able to be used for analysis. Sequences to the *rarb2* and *hoxb5b* MOs were reported previously [8,26]. The total amount of *rarb1* MO injected was 16 ng. The total amount of *rarb2* MO injected was 7 ng. The suboptimal doses used to test genetic interactions were half these concentrations. The amount of *hoxb5b* MO used was 0.25 ng. A cocktail of 4 ng *cyp26a1* MO1 (5'-TCTTATCATCCTTACCTTTTCTTG) and 2 ng *cyp26a1* MO2 (5'-TAAAAATAATACACTACCTGCAAAC) produced a phenotype similar to *gir* mutants [17]. Suboptimal doses used in experiments were 0.9 ng (*cyp26a1* MO1) and 0.45 ng of (*cyp26a1* MO2). For all injection experiments, 3 ng of *p53* MO were used to help suppress non-specific MO-induced cell death [41]. For experiments, the total amount of MO injected was always kept constant by equilibrating the concentrations with Standard Control MO (Gene Tools).

Capped mRNA was made using a Message Machine Kit (Ambion). 150 pg of mRNA was used for over-expression of all mRNAs in all experiments.

Cell culture and luciferase assay

Luciferase reporter assays were performed in HEK 293 cells as previously described [29].

Western blot analysis and ChIP

Western blots were performed as previously described [29]. Mouse monoclonal anti-myc antibody (Covance) was used for both Western blot analysis and ChIP experiments. The dynabeads (Invitrogen) ChIP protocol was adapted from the Dorsky Lab (University of Utah) ZFIN Protocol. qPCR was used to quantify the enrichment of the fragment containing the RARE (DR5) in embryos injected with the *myc-rarb1* mRNA with respect to control uninjected embryos.

Comparison of genomic sequences

The genomic sequence flanking zebrafish *hoxb5b* (-8 to +8 kb) was compared with the corresponding region for *Hoxb5* in mouse using mVista. NHR SCAN was used to identify binding sites for nuclear receptor.

Identification of *rarb1* and RT-PCR

Rarb1 was identified by using BLAST against the zebrafish genome (Ensemble_V7) with the human and mouse RAR α 1 A domains. MacVector was used for sequences alignments. For RT-PCR, primer pairs were designed so that they specifically recognized *rarb1* and *rarb2* (Figure 1B). Primer sequences are available upon request.

Cloning

The full-length coding sequence for *rarb1* was cloned into pCS2p+. The *rarb2*-pCS2p+ construct used for overexpression was reported previously [29]. The myc tagged RAR α 1 was made using the pCS2+MT vector. For *rarb1* and *rarb2* probes, 536 base pairs (bps) of *rarb1* and 443 bps of *rarb2*, which include the 5' untranslated region (UTR) and the specific A domains with no overlap, were cloned (Figure 1B). These fragments were cloned into pGEM-T easy (Promega).

qPCR

Total RNA was isolated from 25 embryos, homogenized in TRIzol (Ambion) and collected using Pure link RNA Micro Kit (Invitrogen). 1 μ g or 0.5 μ g RNA was used for cDNA synthesis using the ThermoScript Reverse Transcriptase kit (Invitrogen). Quantitative real time PCR (qPCR) for *myl7*, *vmhc*, *amhc*, *nkx2.5*, *hand2*, *hoxb5b*, *hoxb5a*, *hoxb6b*, *dhrs3a*, *cyp26a1*, *aldh1a2*, *rdh10a*, *rdh10b*, *rara*, *rarb1*, *rarb2*, *rarya* and *raryb*, *egfp* and *mcherry* was performed using standard PCR conditions in a Bio-Rad CFX PCR machine with Power SYBR Green PCR Master Mix (Applied Biosystems). Expression levels were standardized to *ef1 α* expression and all the data were analyzed using the $2^{-\Delta\Delta CT}$ Livak Method. All experiments were performed in a biological triplicate. Primer sequences are available upon request.

Area and length measurements

Areas of *myl7*, *vmhc* and *amhc* expressing cells were measured using ImageJ and statistical analysis was performed as previously described [26]. Length of *egfp* expression and distance between *hoxb5b* and *egr2b* were measured also using ImageJ and statistical analysis was performed as previously described.

Imaging of zebrafish heart and cell counting

Immunohistochemistry, cell counting and statistical analysis were done as previously described [26].

RA and DEAB treatment

RA and DEAB, treatment of embryos was done as previously described [26,27]. Embryos that have been used for gene expression analysis at 8 somites were treated with 1 μ M DEAB, an Aldh1a2 inhibitor, beginning at 40% epiboly or with 1 μ M RA for 1 hr beginning at 40% epiboly. For analysis of the effects of low concentrations of RA on heart development, embryos were treated with 0.05 μ M RA for 1 hr beginning at 40% epiboly and harvested at 48 hpf. For rescue experiments related to the heart phenotype of RAR α 1 deficient embryos, embryos were treated with 0.025 μ M DEAB beginning at 40% epiboly until 24 hpf. For rescue experiments related to the MHB in RAR α 1+Cyp26a1 deficient embryos, embryos were treated with 0.25 μ M DEAB.

Statistical analysis

To assess whether the means of two groups are statistically different from each other, we applied the Student's *t*-test. A *p* value of <0.05 was considered statistically significant.

Supporting Information

Figure S1 Comparison of RAR α 1 and RAR α 2 expression. (A, D, G, J, M) *rarb1* expression. (B, E, H, K, N) *rarb2* expression. (C, F, I, L, O) *rarb1/2* is a probe that recognizes both isoforms [24]. Arrows in J and L indicate anterior ventral expression. Arrowheads in K and L indicate hindbrain and anterior spinal cord expression. Arrows in M and N indicate differences in the expression of the developing tail. In A–O, all views are lateral. In D–O, dorsal is to the right. (TIF)

Figure S2 RAR α 1 deficient embryos have enlarged hearts at 72 hpf. (A) Control sibling *Tg(-5.1myl7:GFP)^{f2}* embryo. (C) RAR α 1 deficient *Tg(-5.1myl7:GFP)^{f2}* embryo. Arrow in C indicates pericardial edema with enlarged heart. (B, D) Higher magnification images of the fluorescent hearts of the *Tg(-5.1myl7:GFP)^{f2}* control sibling and RAR α 1 deficient *Tg(-5.1myl7:GFP)^{f2}* embryos in A and C. Images are lateral views with dorsal up and anterior right. (TIF)

Figure S3 RAR α 1 deficient embryos have enlarged hearts with increased CM number at 55 hpf. (A, B) Hearts from control sibling and RAR α 1 deficient *Tg(-5.1myl7:DsRed-NLS)^{f2}* embryos at 55 hpf. Images are frontal views. Red indicates ventricle. Green indicates atrium. (C) Mean CM number at 48 hpf. (TIF)

Figure S4 Specificity controls for the translation blocking *rar α 1* MO. (A–C) Control sibling, RAR α 1 deficient, and RAR α 1 deficient+*rar α 1* mRNA injected embryos. Images are lateral views with anterior right at 48 hpf. Red outline indicates ventricles. Green outline indicates atria. Arrow in B indicates edema often found in RAR α 1 deficient embryos, which is not found in RAR α 1 deficient+*rar α 1* mRNA injected embryos (C). (D) qPCR for CM differentiation marker genes at 48 hpf in control sibling, RAR α 1 deficient, RAR α 1 deficient embryos+*rar α 1* mRNA, and RAR α 1 deficient embryos+*kaede* (control) mRNA injected embryos at 48 hpf. Pound sign indicates a statistically significant difference compared to RAR α 1 deficient and RAR α 1 deficient embryos+*kaede* (control) mRNA injected embryos ($p < 0.05$). (TIF)

Figure S5 RARs can directly bind the RA response element (RARE) in the zebrafish *hoxb5b* regulatory region. (A) mVista sequence alignment of mouse *Hoxb5* and zebrafish *hoxb5b* genomic regions. Purple boxes represent exons. Light blue boxes indicates 5' and 3' UTR. Peaks represents levels of sequence identity in a 50 bp window. Purple peaks are conserved regions in exons. Light blue peaks are conserved regions in 5' UTR. Pink peaks are conserved non-coding sequences. Arrow indicates the presence of a RARE in the conserved sequence between 4 kb and 4.5 kb identified previously [42], which we confirmed using the NHR SCAN database. (B) Sequence conservation (red) between mouse and zebrafish DR5 RARE. (C) Western blot for myc-tagged RAR α 1. (D) ChIP from control sibling and *myc-rar α 1* mRNA injected embryos. Negative control *amhc* primers did not detect any enrichment (data not shown). (TIF)

Figure S6 A suboptimal dose of *hoxb5b* MO does not affect CM cell number at 48 hpf. (A, B) Hearts from control sibling and *Hoxb5b* deficient *Tg(-5.1myl7:DsRed-NLS)^{f2}* embryos at 48 hpf. Images are frontal views. Red indicates ventricle. Green indicates atrium. (C) Mean CM number at 48 hpf. (TIF)

Figure S7 RAR α 1 and RAR α 2 function partially redundantly to promote forelimb development. (A–C) Control sibling, RAR α 1 deficient, and RAR α 2 deficient embryos. Images in A–C are dorsal views with anterior to the left. Arrows in B indicate smaller forelimbs. (D) Percentage of control sibling ($n = 20$), RAR α 1 deficient ($n = 20$), and RAR α 2 deficient ($n = 20$) embryos with normal, small or no forelimbs. An optimal dose of the *rar α 1* and *rar α 2* MOs was used for experiments in D. (E) Percentage of embryos with normal, small, or no forelimbs after injection with a suboptimal dose of *rar α 1* MO ($n = 28$), a suboptimal dose of *rar α 2* MO ($n = 26$), and co-injected with suboptimal doses of the *rar α 1* and *rar α 2* MOs ($n = 17$). (F–H) ISH of *tbx5a*, a forelimb marker, in control sibling, RAR α 1 deficient, RAR α 2 deficient embryos. Arrows in F–H indicate *tbx5a* expression the LPM. (I) Areas of the amount of cells expressing the *tbx5a* at 24 hpf. (TIF)

Figure S8 RAR α 2 deficient embryos do not have enlarged hearts. (A, B) Hearts from control sibling and RAR α 2 deficient *Tg(-5.1myl7:DsRed-NLS)* embryos at 48 hpf. Images are frontal

views. Red indicates ventricle. Green indicates atrium. (C) Mean CM number from the hearts of control sibling and RAR α 2 deficient *Tg(-5.1myl7:DsRed-NLS)* embryos at 48 hpf. (D) qPCR for CM marker gene expression in control sibling and RAR α 2 deficient embryos at 48 hpf. We do find a modest decrease in CM number (C) and *myl7* expression (D), which is likely due to a very modest amount of MO-induced toxicity. (TIF)

Figure S9 Patterning of the spinal cord is affected in the RAR α 1+RAR α 2 deficient embryos. (A–D) *Hoxb5a* (spinal cord) and *egr2b* (rhombomeres 3+5) expression in control ($n = 32$), RAR α 1 deficient ($n = 23$), RAR α 2 deficient ($n = 16$), and RAR α 1+RAR α 2 deficient embryos ($n = 19$). (E) Measurements of the distance in arbitrary units (AU) between *hoxb5a* and *egr2b* expression. Expression of *hoxb5a* in the spinal cord is expanded rostrally. The rostral expansion of *hoxb5a* in RAR α 1 deficient embryos trends similarly as RAR α 2 deficient and RAR α 1+RAR α 2 deficient embryos, but it is not statistically significant ($p = 0.06$). (TIF)

Figure S10 RAR α 1 and RAR α 2 function partially redundantly to promote proper heart development. (A–D) Control sibling, RAR α 1 deficient (suboptimal dose), RAR α 2 deficient (suboptimal dose), and RAR α 1+RAR α 2 (suboptimal doses) deficient embryos at the 72 hpf. Arrow in D indicates pericardial edema and the enlarged heart. (TIF)

Figure S11 *Rar α 1* and *rar α 2* mRNA overexpression do not significantly affect RA responsive genes. (A–I) ISH for the RA responsive genes *cyp26a1*, *dhrs3a*, and *hoxb5b* at 8 s. (A, D, G) Control sibling, (B, E, H) *rar α 1* mRNA, and (C, F, I) *rar α 2* mRNA injected embryos. Injection of either *rar α* mRNA did not inhibit RA responsive gene expression. Images in A–C are lateral views with anterior up and dorsal right. Images in D–I are dorsal views with anterior up. (J) qPCR for RA responsive genes *cyp26a1*, *hoxb5a*, *hoxb8b*, *dhrs3a* at 8 s. (K) Mean CM number from control sibling, *rar α 1* mRNA, and *rar α 2* mRNA injected *Tg(-5.1myl7:DsRed-NLS)^{f2}* embryos. (L) Transfection of HEK 293 cells with DNA for the zebrafish *rar α 1* and *rar α 2* and pGL3-12XRARE-*ef1 α :renilla luciferase* vector with and without RA treatment. Fold difference in luminescence is indicated in arbitrary units (AU) and reflects the ratio of renilla luciferase (RL) to firefly (FL) luciferase. (TIF)

Figure S12 *Rar* expression in RAR α 1, RAR α 2, or RAR α 1+2 deficient embryos. (A, B) ISH for *rar α 1* in RAR α 2 deficient embryos. (C, D) ISH for *rar α 2* in RAR α 1 deficient embryos. (E, F) ISH for *rar α 2* in RAR α 1+2 deficient embryos. (G, H) ISH for *rarg α* in RAR α 1+2 deficient embryos. (I, L) ISH for *rarg β* in RAR α 1+2 deficient embryos. *rar* expression is often expanded in the tailbud region of embryos deficient for the other RAR homologs, while additional regions also appear to have increased or low levels of ectopic expression. All views are lateral with dorsal right at 8 s. Arrows in A–H indicate distance of expression in the tail. Arrowheads in F, H, L indicate regions of increased or ectopic expression. (TIF)

Figure S13 *Rdh10a* expression in RAR α 1+2 deficient embryos. (A–D) ISH for *rdh10a* in RAR α 1+2 deficient embryos at 8 somites. (A, B) Lateral views with dorsal right. (C, D) Dorsal views with anterior up. Brackets indicate expansion of *rdh10a* in the ALPM. Arrow indicates increased expression in the somites. (TIF)

Figure S14 Characterization of *cyp26a1* splice-blocking MOs used in experiments. (A) Schematic of the *cyp26a1* locus and the intron-exon boundaries targeted by the different *cyp26a1* MOs. Blue bar indicates MO1. Red bar indicates MO2. MO1 primarily causes usage of two in-frame cryptic splice sites. Dashed lines indicate the alternate introns cause by the cryptic splices induced from MO1. MO2 causes the introduction of a premature stop codon (red X). (B) RT-PCR for the WT *cyp26a1* transcripts and alternate transcripts induced from the different MOs. U and L indicate bands depicted in A. (C) Control sibling embryo. (D) Embryos injected with cocktail of *cyp26a1* MO1+2. Co-injection of *cyp26a1* MO1 and MO2 causes a phenotype equivalent to or stronger than the *cyp26a1/giraffe (gir)* mutant (E). Injection of the individual MOs causes the phenotypes consistent with *cyp26a1* loss of function at low frequency (data not shown). A suboptimal dose of the *cyp26a1* MO cocktail was used for functional interaction experiments with RAR α b1 (Figure 4). Arrows in D and E indicate shortened tail. Views in C–E are lateral with anterior right. (TIF)

Figure S15 Characterization of the novel transgenic RA sensorline. (A) Schematic of the RAR domains and the Gal4 DNA binding domain (GDBD)/RAR α b ligand binding domain (RLBD) fusion protein. Grey indicates the GDBD. Yellow indicates the RLBD. D is a linker domain and F is a domain with unknown function (as in Figure 1). (B, C) Schematics

representing the GDBD-RLBD fusion acting on the *Gal4-UAS:EGFP* transgene. The GDBD-RLBD is expressed under the β -actin promoter. (B) In the absence of RA, *egfp* is not expressed. (C) In the presence of RA (red triangles), the GDBD-RLBD is able to promote *egfp* (UAS responsive gene) transcription. (D–G) *Tg(β -actin:GDBD-RLB);Tg(UAS:nfsB-mcherry)* embryos are responsive to RA treatment. ISH for *mcherry*. Equivalent results were found when the *Tg(β -actin:GDBD-RLB)* line was crossed to *Tg(UAS:EGFP)* fish (data not shown) as were used for experiments in Figure 5. More detailed characterization of the stable transgenic RA sensor lines is reported in [31]. (D, E) Lateral views with dorsal right. (F, G) Dorsal views. In images D–G anterior is up.

(TIF)

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Author Contributions

Conceived and designed the experiments: ED JSW. Performed the experiments: ED ABR JLA AM JSW. Analyzed the data: ED ABR AM JSW. Contributed reagents/materials/analysis tools: ED ABR JLA AM JSW. Wrote the paper: ED JSW.

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