Transcriptomic Analysis of Murine Embryos Lacking Endogenous Retinoic Acid Signaling

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Abstract

Retinoic acid (RA), an active derivative of the liposoluble vitamin A (retinol), acts as an important signaling molecule during embryonic development, regulating phenomena as diverse as anterior-posterior axial patterning, forebrain and optic vesicle development, specification of hindbrain rhombomeres, pharyngeal arches and second heart field, somitogenesis, and differentiation of spinal cord neurons. This small molecule directly triggers gene activation by binding to nuclear receptors (RARs), switching them from potential repressors to transcriptional activators. The repertoire of RA-regulated genes in embryonic tissues is poorly characterized. We performed a comparative analysis of the transcriptomes of murine wild-type and Retinaldehyde Dehydrogenase 2 null-mutant (Raldh2−/−) embryos — unable to synthesize RA from maternally-derived retinol — using Affymetrix DNA microarrays. Transcriptomic changes were analyzed in two embryonic regions: anterior tissues including forebrain and optic vesicle, and posterior (trunk) tissues, at early stages preceding the appearance of overt phenotypic abnormalities. Several genes expected to be downregulated under RA deficiency appeared in the transcriptome data (e.g. Emx2, Foxq1 anteriorly, Cdx1, Hoxa1, Rab2 posteriorly), whereas reverse-transcriptase-PCR and in situ hybridization performed for additional selected genes validated the changes identified through microarray analysis. Altogether, the affected genes belonged to numerous molecular pathways and cellular/organismal functions, demonstrating the pleiotropic nature of RA-dependent events. In both tissue samples, genes upregulated were more numerous than those downregulated, probably due to feedback regulatory loops. Bioinformatic analyses highlighted groups (clusters) of genes displaying similar behaviors in mutant tissues, and biological functions most significantly affected (e.g. mTOR, VEGF, ILK signaling in forebrain tissues; pyrimidine and purine metabolism, calcium signaling, one carbon metabolism in posterior tissues). Overall, these data give an overview of the gene expression changes resulting from embryonic RA deficiency, and provide new candidate genes and pathways that may help understanding retinoid-dependent molecular events.

Citation: Paschaki M, Schneider C, Rhinn M, Thibault-Carpentier C, Dembélé D, et al. (2013) Transcriptomic Analysis of Murine Embryos Lacking Endogenous Retinoic Acid Signaling. PLoS ONE 8(4): e62274. doi:10.1371/journal.pone.0062274

Editor: Michael Schubert, Laboratoire de Biologie du Développement de Villefranche-sur-Mer, France

Received November 21, 2012; Accepted March 19, 2013; Published April 24, 2013

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Funding: This work was supported by grants from the Agence Nationale de la Recherche (ANR Neurosciences 2007, ANR Blanc 2011) and the Fondation pour la Recherche Médicale (Equipe FRM 2007), and by institutional funds from the Centre National de la Recherche Scientifique (CNRS) and Institut National de la Santé et de la Recherche Médicale (INSERM). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

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Introduction

Retinoic acid (RA), an active derivative of the liposoluble vitamin A, is an endogenous signaling molecule involved in many biological processes in vertebrates. This small hydrophobic compound is the ligand for a subfamily of nuclear receptors, the retinoic acid receptors (RARs), switching them from potential repressors to transcriptional activators. The repertoire of RA-regulated genes in embryonic tissues is poorly characterized. We performed a comparative analysis of the transcriptomes of murine wild-type and Retinaldehyde Dehydrogenase 2 null-mutant (Raldh2−/−) embryos — unable to synthesize RA from maternally-derived retinol — using Affymetrix DNA microarrays. Transcriptomic changes were analyzed in two embryonic regions: anterior tissues including forebrain and optic vesicle, and posterior (trunk) tissues, at early stages preceding the appearance of overt phenotypic abnormalities. Several genes expected to be downregulated under RA deficiency appeared in the transcriptome data (e.g. Emx2, Foxq1 anteriorly, Cdx1, Hoxa1, Rab2 posteriorly), whereas reverse-transcriptase-PCR and in situ hybridization performed for additional selected genes validated the changes identified through microarray analysis. Altogether, the affected genes belonged to numerous molecular pathways and cellular/organismal functions, demonstrating the pleiotropic nature of RA-dependent events. In both tissue samples, genes upregulated were more numerous than those downregulated, probably due to feedback regulatory loops. Bioinformatic analyses highlighted groups (clusters) of genes displaying similar behaviors in mutant tissues, and biological functions most significantly affected (e.g. mTOR, VEGF, ILK signaling in forebrain tissues; pyrimidine and purine metabolism, calcium signaling, one carbon metabolism in posterior tissues). Overall, these data give an overview of the gene expression changes resulting from embryonic RA deficiency, and provide new candidate genes and pathways that may help understanding retinoid-dependent molecular events.

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spatial RA distributions are critical for the establishment and patterning of the hindbrain segments (the rhombomeres), whereas more caudally, RA has been implicated in the regulation of the symmetrical progression of somitogenesis, and in early neurogenic events in the neural tube (the prospective spinal cord) [8,9,10,11] (refs. [12,13,14] for reviews). Other early embryonic functions have been reported for development of the segmented pharyngeal apparatus (the branchial arches), and in cell populations giving rise to the heart [15,16,17,18].

The distribution of RA among cell populations is controlled by specific enzymatic pathways. Embryos from placental species obtain vitamin A in the form of retinol transferred from the maternal circulation through the embryonic yolk sac – and eventually the placenta [19,20]. Oviparous species store retinol and/or carotenoids in the egg yolk. Retinol reaching the embryo can be oxidized to retinaldehyde by two classes of enzymes (alcohol and retinol dehydrogenases: ADHs, RDHs), whereas β-carotenoids are cleaved by β-carotene-15,15′ oxygenases (BCOs) (refs. [21,22,23,24] for reviews). Critical for the production of RA is the presence of specific retinaldehyde dehydrogenases (RALDHs). Three such enzymes have been characterized, all of which being expressed according to distinct temporal and spatial (tissue-specific) patterns during development. Interestingly, there is a temporal window during which RALDH2 is the only RALDH to be expressed in the early embryo, and all the functions mentioned above have indeed been ascribed to this enzyme. This phase goes from gastrulation to early somitic stages, during which Raldh2 gene expression is first induced in the embryonic node and newly formed mesoderm, and then persists at specific anterior-posterior axial levels in several mesodermal derivatives [6,25]. Analysis of murine Raldh2−/− null mutants harboring a RA-activatable reporter transgene has demonstrated the absence of any detectable RA activity throughout the embryo at these early stages [26]. Raldh2 expression is transiently seen at early somite stages (E8.0–E8.5 in the mouse) in head tissues, including the anteriormost neuroectoderm destined to form the anterior forebrain and optic vesicles [27], and knockout mutation of Raldh2 affects morphogenesis of these structures [28,29]. The second next Raldh gene to be expressed is Raldh3, whose transcripts first appear by E8.5 in head structures (prospective nasal region, eye, and otocyst), and at trunk levels in the mesonephric duct (embryonic kidney) [30,31].

Although Raldh2−/− mouse mutants have been used in a large set of studies for unravelling the consequences of a lack of RA signaling in early developmental processes, there is still a poor understanding of the molecular events leading to the realization of the RA-deficiency phenotypes. Genes abnormally expressed in the Raldh2−/− embryos were usually identified through in situ hybridization studies using molecular markers of the affected structures, and there has been no attempt to globally analyze gene expression changes at the transcriptomic level. As this mutant is a good model for analyzing embryos developing under RA deficiency, we decided to perform an Affymetrix DNA microarray study to compare the transcriptomes of wild-type and Raldh2−/− mutants: B, D) at the stages chosen for analysis. The embryos are not from the experimental series, but were collected from litters harboring a retinoic acid-sensitive (RARE-hsp68-lacZ) reporter transgene and processed for X-gal analysis, thereby illustrating the absence of detectable RA activity in the mutants. The collected samples were the anterior head region of 14 somite stage embryos (X-gal-labelled in control embryos, section planes illustrated by arrows: A, B) and the posterior region (X-gal-labelled in control embryos, section plane illustrated by arrows: C, D) of 4 somite stage embryos. (E, F) Principal component analysis illustration of ANT and POST data sets. Wild-type samples are depicted by blue dots, and Raldh2−/− samples by red dots. For anterior tissues, wild-type and Raldh2−/− samples segregate in two distinct groups. In the case of posterior tissues, one Raldh2−/− sample (#046) did not cluster with the others, and was excluded from subsequent analyses. (G) VENN diagrams summarizing the numbers of genes showing statistically significant different expression levels, with a fold change of $±1.2$ (upper numbers) or $±1.5$ (lower, bold numbers). The numbers of genes exhibiting a “contradictory” behavior (upregulated in one type of samples, and downregulated in the other, as pointed out by the arrows) are also given.

doi:10.1371/journal.pone.0062274.g001

Results and Discussion

Experimental design

Two sets of samples were used for analyzing transcriptome changes in Raldh2−/− embryos. The rostral part of the head (including the anterior forebrain, optic vesicles, and overlying tissues), was collected from wild-type and mutant embryos at the 14 somite stage. This stage was chosen according to the expression
profiles of Raldh genes in head tissues of wild-type embryos: it shortly follows the phase of peak Raldh2 mRNA expression at 8-10 somites [27], while RALDH2 protein is still detectable [29], whereas Raldh3 mRNA expression, first detected at the 10-12 somite stage, has not yet intensified. It is unlikely that RALDH3 significantly participates to RA synthesis in the head at the stage of analysis, and – as described in a previous study [29] – we confirmed by analyzing Raldh2−/− mutants carrying the RARE-hsp68-lacZ transgene [32], that there was no detectable activity of the lacZ reporter at the 14 somite stage (Fig. 1A, B). The tissues collected for analysis included the optic cup and the forebrain neuroepithelium located more rostrally (prospective telencephalon), as well as the overlying tissues (frontonasal region including the olfactory placode). These samples are referred to thereafter as “anterior” (ANT) tissues.

The posterior tissues were analyzed at the 4 somite stage, and samples were collected from a transverse section plane excluding all tissues from the level of the first branchial arch (Fig. 1C, D). At this stage RA has numerous documented effects (see Introduction), however Raldh2−/− embryos do not yet exhibit morphological deficiencies (see Fig. 1D) – but they can be recognized by trained eyes by the presence of smaller somites and an abnormally shaped heart tube. As expected from previous work [26], there was no detectable activity of the RARE-hsp68-lacZ transgene in the mutant embryos at this stage (Fig. 1D). These samples will be referred to as “posterior” (POST).

Overview of differentially expressed genes

Using as a threshold the p-values defined above by Partek analysis, 7177 genes were differentially expressed in wild-type vs. mutant ANT tissue samples (3437 upregulated, 3740 downregulated in mutants; p≤0.05). When POST samples were analyzed by omitting the dubious Raldh2−/− sample (#046, Fig. 1F), 4133 genes were found to be differentially expressed (1834 upregulated, 2299 downregulated in mutants; p≤0.03). When analysis was performed including sample #046, 3120 genes were significantly upregulated (24.5% gene loss), and only 1610 genes appeared downregulated (30% gene loss). Thus, as predicted by PCA, inclusion of sample #046 biased the statistical tests, and data mining was performed with this sample excluded.

In both the ANT and POST microarray experiments, the large majority of genes were affected by fold changes (FC) lesser than 2 fold (only 104 genes with FC >2, and 30 genes with FC <−2, in Raldh2−/− vs. wild-type ANT samples; 42 genes with FC >2 and 82 genes with FC <−2, in Raldh2−/− vs. wild-type POST samples). VENN diagrams in Fig. 1G show gene numbers obtained for two cut-offs at ±1.2 FC (upper lines) and ±1.5 FC (lower lines, numbers in bold). The 1.2 FC cut-off yielded comparable numbers of genes up- or downregulated in each experiment, with lower numbers of downregulated genes. Furthermore, there were almost 500 genes upregulated in both the ANT and POST Raldh2−/− tissue samples, and nearly as many genes downregulated in both Raldh2−/− tissue samples. These lists were reduced to 102 genes up-regulated in common, and 14 genes down-regulated in common, at ±1.5 FC. Remarkably, the numbers of “contradictory” genes (being upregulated in one tissue and downregulated in the other) was extremely low (0 and 2 contradictory genes above 1.5 FC; Fig. 1G). Table 1 provides a list of the most severely downregulated genes, and details on contradictory genes are provided as supplementary online information (Table S1). A graphic illustration of the differentially expressed genes is also shown (Fig. 2).

Down-regulated genes include known RA-dependent genes

As RARs behave as transcriptional activators in the presence of their ligand, down-regulation of retinoid target genes is expected to occur in tissues deficient in RA synthesis. We analyzed the lists of downregulated genes to see whether genes already known to be downregulated in the Raldh2 mutant, or more generally reported in the literature as being RA-dependent, were detected in our microarray experiments. Several genes shown in previous studies to be downregulated in Raldh2−/− embryos (usually including whole-mount in situ hybridization) [6,26,33,34,35] were also found downregulated in the POST microarray experiment. These include Rarb, Ptx6, Hoxa1 (all <−2), Meox1 (−1.68), and Cdx1 (−1.40). Additional RA targets were reduced in the POST experiment, although excluded from the filtered list due to their p-values, including Neurog2 (−1.24) [36], Olg2 (−1.78) and Tbx5 (−1.35) [18]. Stra7, encoding a membrane protein interacting with retinol-binding protein (RBP) and originally characterized as a RA-induced gene in P19 embryonal carcinoma cells [37,38] and Dhrs3 (also known as RetSDR1), encoding a retinaldehyde dehydrogenase/reductase, were both reduced in the caudal region. Downregulation of Dhrs3 may be due to a feedback mechanism caused by retinaldehyde accumulation in embryonic tissues [39]. The Cspa2 gene (encoding cellular RA-binding protein-II) was downregulated in both posterior and anterior tissues, indicating that the retinoic acid-response elements (RAREs) identified in the Cspa2 gene promoter [40] are functional targets. The Nkx3-1 and Nkx3-2 genes were both
reduced in posterior tissue samples. *Nkx3-1* is a RA-inducible target in prostate cancer cells [41].

Several genes previously shown be reduced in the forebrain of *Raldh2*−/− embryos [29] were also downregulated in our ANT microarray data, for instance *Dll1* (−1.31), *Fog1* (−1.42) and *Emx2* (−1.56). Thus, a good overall correlation between our transcriptomic data and previous analyses of *Raldh2*−/− embryos (typically by non quantitative in situ hybridization) confirms the validity and reproducibility of these studies.

**Microarray analysis provides new candidate genes for the RA-deficiency phenotypes**

When considering the list of genes exhibiting the highest changes in expression levels in *Raldh2*−/− embryos (Table 1), some interesting candidates emerged as being potentially involved in the RA-deficiency phenotype. The most highly downregulated genes in anterior tissues encode Id4 (inhibitor of DNA binding 4), a regulator of neural stem cell proliferation and differentiation [42], in anterior tissues encode Id4 (inhibitor of DNA binding 4), a regulator of neural stem cell proliferation and differentiation [42], the signal transducer and activator of transcription Stat4, and the regulator of neural stem cell proliferation and differentiation [42], in anterior tissues encode Id4 (inhibitor of DNA binding 4), a regulator of neural stem cell proliferation and differentiation [42].

Among the most severely downregulated genes are also the Ephrin receptor gene *EphA3*, the neuropilin/tolloid-like gene *Neto2*, and the gene encoding fatty acid-binding protein 7 (Fabp7 or brain-type Fabp). The *N2r1* gene, encoding the orphan nuclear receptor Tfrx1, was also strongly downregulated in anterior tissues. A previous study pointed to Tfrx1 as being involved in the regulation of the *Rarb* gene in the eye [43]. The SRY-box genes *Sox1* (−1.66) and, to a lesser extent, *Sox3* (−1.26), were also downregulated according to the ANT microarray experiment.

Retinoids orchestrate the differentiation and patterning of the posterior neural tube (the developing spinal cord) [7,13], accompanied by induction of a posterior transcriptional signature previously analyzed in the spinal cord at an early fetal stage [44]. Our analysis of posterior embryonic tissues (see Table 1) confirmed many known caudal neuronal targets, including *Pax6*, *Hoxa1*, *Dlx1*, a homeobox gene involved in spinal cord interneuron specification [45] and *Lhx1*, a LIM-homeodomain protein regulating gastrulation, renal organogenesis, and other embryonic functions [46]. Of note, the *Lhx1* homologue *Lhx2* was downregulated in both ANT (−1.60) and POST (−1.45) tissues. Novel downregulated genes included *Shp5*, encoding a secreted antagonist of Wnt signaling [47], and *Netn* (*Nephrocyst*), encoding an inhibitor of TGFβ signaling [48]. Other significant reduced targets include *Dmnt2* (doublesex and mab-3 related transcription factor 2), involved in somitogenesis and early myogenesis [49], *Myc* (a regulator of early myogenic lineage decisions), *Metn* (encoding Meteorin, a secreted protein controlling neurogenesis and angiogenesis in glial precursors [50]), *Tlca* (encoding a Netrin receptor involved in spinal cord development [51]), *Tif5* (encoding the basic helix-loop-helix protein Paraxis involved in pre-somatic mesoderm differentiation [52]), and *Rgs6* (encoding a winged helix transcription factor required for pancreatic development [53]).

While a deficiency in embryonic RA should primarily result in lack of activation of RA/RAR inducible genes, feedback mechanisms may lead to upregulation of genes which may also be relevant for the resulting phenotypes. Among the highest upregulated genes were two genes coding for basic helix-loop-helix proteins, *Bhlhe40* and *Bhlhe41* also known as *Shap2* and *Shap1* [54]. *Bhlhe40* was first reported under the name of *Stox13*, as a RA-inducible gene promoting neuronal differentiation in P19 cells [55]. It was therefore unexpected to see it upregulated in vivo in *Raldh2*−/− embryonic tissues. For this reason we included *Bhlhe40* among the set of genes for which additional analyses (quantitative RT-PCR and in situ hybridization) were performed to confirm the expression changes (see below). Other transcription factor-coding genes were upregulated in *Raldh2*−/− samples, namely the *Hairy/Enhancer-of-Split*-related gene *Hsy2*, the winged-helix protein-coding gene *FoxO3*, the zinc finger protein-coding gene *Egr1* (formerly known as *Krox24*), and the orphan nuclear receptor *ROR* gene (*Rora*). Other upregulated genes included the *Ras* proto-oncogene coding for the R-Ras GTPase, the *Ccn2* gene coding for cyclin G2 – which is regulated at the mRNA level by *FoxO3* [56], and the *Vldh* (very low density lipoprotein receptor) gene, the latter protein also acting as a Reelin receptor during development [57]. Genes encoding ligands or modulators of specific signaling pathways (e.g. *Efn3*, *Fgfl1*, *Igfbp1*, *Vgfa*) were also present in the list. A detailed analysis of the main genetic and
| Symbol | Name | Function |
|--------|------|----------|
| Id4    | Inhibitor of DNA binding 4 | Regulation of transcription and cell proliferation, brain, hippocampus development |
| Stat4  | Signal transducer and activator of transcription 4 | Regulation of transcription and cell proliferation, signal transduction |
| Taflb  | Taflb, TBP-associated factor | Regulation of transcription |
| Mfsd2  | Major facilitator superfamily domain containing 7B | Unknown |
| Car3   | Carbonic anhydrase 3 | Lyase activity |
| EphA3  | Eph receptor A3 | Transmembrane receptor, protein kinase |
| Rhag   | Rhesus blood group-associated A glycoprotein | Ammonium transmembrane transporter |
| Syl11  | Synaptotagmin XI | Calcium transmembrane transporter |
| Prx21  | Protein tyrosine phosphatase, receptor type Z, polypeptide 1 | Axonogenesis |
| Cpa2   | Carboxypeptidase A2 | Metalloendopeptidase |
| Neto2  | Neuropilin (NRP) and tolloid (TLL)-like 2 | Membrane receptor |
| Gm9983 | Predicted gene 9983 | Unknown |
| Mfsd7b | Major facilitator superfamily domain containing 7B | Head, digits, and spleen development |
| Fabp7  | Fatty acid binding protein 7, brain | Lipid transport, forebrain neurogenesis |
| Slfn9  | Schlafen 9 | Unknown |
| Jakmip2| Janus kinase and microtubule interacting protein 2 | Kinase, microtubule interacting protein |
| Slc4a1 | Solute carrier family 4 (anion exchanger), member 1 | Anion exchanger transport |
| Ntr2e1 | Nuclear receptor 2E1 (Tlx) | Regulation of transcription, brain and eye development |
| Id1    | Isopentenyl-diphosphate delta isomerase | Steroid and carotenoid biosynthesis |
| Lonf2  | LON peptidase N-terminal domain and ring finger 2 | Unknown |

**DOWN in POSTERIOR TISSUES**

| Symbol | Name | Function |
|--------|------|----------|
| Arg1   | Arginase | Arginase activity |
| 1700011H14Rik | Unknown | Unknown |
| Rarb   | Retinoic acid receptor beta | Transcriptional regulation, embryogenesis, neurogenesis |
| Nepn   | Nephrocan | Negative regulation of transcription and of TGF-β receptor signaling |
| Hnf1b  | HNF1 homeobox B | Regulation of transcription, inner cell mass differentiation, anterior-posterior patterning, hindbrain development |
| Dlx1   | Developing brain homeobox 1 | Regulation of transcription, spinal cord interneuron specification |
| Rfx6   | Regulatory factor X, 6 | Regulation of transcription, pancreatic development |
| Pax6   | Paired box gene 6 | Regulation of transcription, brain, eye, spinal cord development, neurogenesis |
| Hoxa1  | Homeobox gene A1 | Regulation of transcription, hindbrain development, anterior-posterior patterning, inner ear morphogenesis |
| Ntx3   | Nuclear RNA export factor 3 | mRNA export |
| Gm10785| Unknown | Unknown |
| Unc5a  | Unc-5 (C. elegans) homolog A | Membrane receptor, apoptosis, signal transduction |
| Maob   | Monoamine oxidase B | Oxidoreductase |
| Lhx1   | LIM homeobox protein 1 | Regulation of transcription, ectoderm/endoderm formation, pattern specification, nervous system and urogenital development |
| Slc16a12| Solute carrier family 16 (monocarboxylic acid transporters), member 12 | Sympporter activity |
| Olig2  | Oligodendrocyte transcription factor 2 | Regulation of transcription, oligodendrocyte specification |
| Fgfr4  | Fibroblast growth factor receptor 4 | Membrane receptor, tyrosine kinase |
| Sfrp5  | Secreted frizzled-related sequence protein 5 | Extracellular modulator of Wnt signaling, foregut endoderm and mesoderm development |
| Cpa1   | Carboxypeptidase A1 | Metalloendopeptidase, proteolysis |
| Dhrs3  | Dehydrogenase/reductase (SDR family) 3 | Oxidoreductase |
Genes listed alphabetically, fold change in expression
Genes listed by decreasing fold change in expression, ranging from
Genes listed by decreasing fold change in expression, ranging from
The table distinguishes the “top” downregulated genes in anterior or posterior tissues, and the genes affected in both types of samples.
doi:10.1371/journal.pone.0062274.t001

Table 1. Cont.

DOWN in ANTERIOR and POSTERIOR TISSUES (3)
Symbol Name Function

| Symbol | Name | Function |
|---------|------|----------|
| Apba2  | Amyloid beta (A4) precursor protein-binding, family A, member 2 | Neuronal adaptor protein, synaptic transmission |
| Ap1s3  | Adaptor-related protein complex AP-1, sigma 3 | Intracellular protein transport, endocytosis |
| Ccnb1p1| Cyclin B1 interacting protein 1 | Blasto cyst formation, organ growth, meiosis and spermatid development, apoptosis |
| Ccne1  | Cyclin E1 | Cyclin-dependent protein kinase, cell cycle control |
| Con3   | Casein kappa | Oocyte meiosis, lactation |
| Uolv2  | Elongation of very long fatty acids-like 2 | Lipid biosynthesis |
| Ktbd8  | Kelch repeat and BTB (POZ) domain containing 8 | Unknown |
| Pmn2   | Phosphomannomutase 2 | Mannose biosynthesis |
| Ptprz1  | Protein tyrosine phosphatase, receptor type Z | Extracellular matrix protein, hydrolase, neurogenesis |
| Rpl12  | Ribosomal RNA processing 12 (S. cerevisiae) homolog | Unknown |
| Snhg1  | Small nuclear RNA host gene | Unknown |
| Tmem56 | Transmembrane protein 56 | Unknown |
| Tmmi61a| tRNA methyltransferase 61 homolog A | tRNA methyltransferase activity |
| Pinx1  | PIN2/TERF1 interacting, telomerase inhibitor 1 | Telomere maintenance, nucleic acid and protein binding |

(1) Genes listed by decreasing fold change in expression, ranging from –3.3 to –1.9.
(2) Genes listed by decreasing fold change in expression, ranging from –4.5 to –1.7.
(3) Genes listed alphabetically, fold change in expression < –1.5 in both the ANT and POST microarrays.

Retinoic acid is commonly used as an inducer of neuronal differentiation in various tissue culture cell lines. Remarkably, retinoids can differentiate embryonic stem (ES) cell-derived embryoid bodies to a relatively pure stem cell lineage known as radial glia [61]. Radial glial cells are found at the earliest stages of CNS development and play an important role in cortical expansion. An in vivo role of retinoids in promoting this lineage, though, has never been shown. Two players in the radial glia lineage, Pax6 and Fabp7, are both reduced in ANT Raldh22−/− tissues. Pax6 maintains the proliferative capacity and developmental potential of radial glia stem cells [62], whereas Fabp7 (also known as Brain Lipid Binding Protein, BLBP) is a structural marker also required for the establishment of the radial glia fiber system in the developing brain [63].

Clustering analysis of gene expression changes in Raldh22−/− embryos

To obtain a more global view of the gene expression changes detected in the Raldh22−/− embryos, we performed clustering analysis using the Cluster 3 software [64]. An overview of the entire clustering analysis is provided as online supporting information in the form of a heat map (Figure S1), and selected gene clusters are shown in Figure 3. A green-to-red scale (Fig. 3, bottom) illustrates the gene expression changes between samples, from lowest (bright green) to highest (bright red) relative levels. The clustering analysis further validated our experimental samples, as WT and mutant (KO) samples segregated into distinct clusters, with no co-segregation of any WT and KO sample (Fig. 3A).

regulatory pathways emerging from the transcriptomic analysis of RA-deficient embryos is presented below.

Retinoic targets regulating stem cell lineages

Our transcriptomic analysis has uncovered some novel RA targets regulating neuronal stem cell lineages, and whose deficiency might explain forebrain abnormalities in Raldh22−/− embryos. Id4, Tlx, and Sox1 are all reduced in ANT Raldh22−/− tissues. The mutant embryos exhibit dramatic reductions in telencephalon neuroepithelium growth, a defect previously proposed to be due to reduced fibroblast growth factor (FGF) and Sonic hedgehog signaling [29]. Retinoic regulation of gene targets in stem cells could also stunt cerebral growth. Interestingly, Id4 regulates lateral expansion of the proliferative zone in the developing cortex [42] by facilitating self-renewal and proliferation of neural stem cells [58]. Tlx maintains neural stem cells in an undifferentiated, proliferative state [59], whereas Sox1-expression marks activated neural progenitors. Sox1 also functions to maintain cortical progenitors in an undifferentiated state by suppressing cell cycle exit to neurogenesis [60]. Collectively, Id4, Tlx, and Sox1 reductions may be sufficient to affect forebrain development by hindering neural stem cell growth. Neural stem cells may consequently undergo premature differentiation, explaining why we observe increased levels of Sox10 [29] and Bhlhe40 in Raldh22−/− forebrains. Reduced FGF signaling and cyclin D levels [29] could be a consequence, and not the cause, of reduced progenitor growth. In another study we have shown that Raldh2 loss of function reduces expansion and differentiation fetal spinal cord-derived neural stem cells [44]. Our ongoing work using neurosphere assays indicates that neural stem cells are similarly affected in the forebrain of 14 somite-stage Raldh22−/− embryos (M.P. et al, unpublished data).
Such analysis allows us to pinpoint genes exhibiting similar behaviors (in terms of variation in expression levels) among the analyzed samples. As a first example, the Id4 and protocadherin 19 (Pcdh19) genes clustered with Six2, being selectively downregulated in anterior tissue samples (Fig. 3B). Interestingly also, the Foxg1 and Emx2 genes – both described as downregulated in the forebrain of Raldh2−/− embryos [29] – were found in the same cluster (Fig. 3C). This cluster also included a Wnt gene (Wnt7b), the LIM/homeobox gene Lhx2, and the nuclear receptor geneNr2e1. Another interesting cluster contained the Pax6 and Crabp2 genes (Fig. 3D). Unlike the former ones, genes from this cluster were downregulated in both anterior and posterior Raldh2−/− tissues samples, while their expression levels were higher in anterior tissues (both in WT and KO situations). Such a behavior is not surprising, as Pax6 and Crabp2 are highly expressed in most of the tissue (forebrain and optic vesicle) constituting the ANT samples. The cluster also included Sall2 (a drosophila Spalt homologue), Unc119b, Mkrn3 (Makorin), and Metrn (Meteorin). Another cluster grouped the Msx2 homeobox gene, a Snail homologue (Snai2), a Dickkopf homologue (Dkk3), an Ikaros family gene (Ikzf4), and the Map kinase gene Map4k2, all being relatively mildly downregulated in ANT samples (Fig. 3F).

Genes from the cluster containing Foxo3 and the IGF receptor gene Igf1r, on the other hand, were upregulated in both the ANT

Figure 3. Details of the gene expression profiles obtained after hierarchical clustering of the experimental samples by relative gene expression level analysis. Eight clusters (panels B–I) have been extracted from the overall clustering analysis (available as additional online information, Figure S1), showing differential gene expression behaviors according to the experimental samples (ANT, RNA from anterior tissues; POST, RNA from posterior tissues; WT, wild-type embryos; KO, Raldh2−/− embryos). Gene expression profiles are illustrated as a heat map (green: weak expression; red: strong expression – see scale below). The expression profile of the Rab gene is also shown (panel J), which did not cluster with any other gene. This analysis further validated the experimental samples, as WT and KO samples segregated into fully distinct clusters, both for the ANT and POST tissue samples (panel A above). doi:10.1371/journal.pone.0062274.g003

Figure 4. Microarray validation by quantitative real-time RT-PCR (Q-PCR). Genes were selected according to their different expression profiles as detected by microarray analysis of wild-type (WT) and Raldh2−/− embryos. Among these, one gene (Pax6) was chosen as a “control”, as its expression had already been shown to be affected in Raldh2−/− embryos (see main text for details and interpretation of the data). Expression levels are expressed as values normalized with respect to Gapdh (a housekeeping gene) mRNA levels (WT samples, blue bars; Raldh2−/− samples, red bars). Data (mean ± SEM) were analyzed with Student t-test; ***, p<0.001; **, p<0.01; *, p<0.05. doi:10.1371/journal.pone.0062274.g004
and POST Raldh2−/− samples, although globally they were expressed at higher levels posteriorly (Fig. 3G). This 14 gene cluster also contained the chordin-like gene Cnml, the latent 
TGFβ-binding protein gene Lhbp1, the zinc finger protein genes Zksam2 and Zngm6, as well as 3 unannotated Riken gene transcripts. 

Two examples of genes exhibiting selective downregulation in the POST Raldh2−/− tissue samples are shown. In the first one, the RARE-containing gene Cdx1 clusters with Lhx1, Sfrp5, the nephrocan (Nepo) gene, and the gene encoding the FGF pathway inhibitor Sprouty3 (Spy3) [Fig. 3H]. The second one includes Cil1, encoding a transcriptional effector of Hedgehog signaling shown to be downregulated in Raldh2−/− embryos [6]. This gene clusters with Sost6 and several other potentially interesting genes, including two NKx genes (NKx2-2 and 2-9), the Frizzled homologue Fzd4, the forkhead protein-coding gene Foxp4, Ripply3 (encoding a Tbx1 repressor), and Unc5a (encoding a Netrin receptor).

Finally, the gene encoding RARβ (Rarb), a direct RA-inducible gene containing a functional RARE [65] was not included in any gene cluster (Fig. 3I). This behavior could be linked to its weak expression in all anterior tissue samples, whereas downregulation was clearly seen in posterior samples.

Selected gene analysis by quantitative RT-PCR and in situ hybridization

Some genes were selected to perform real-time quantitative reverse transcriptase-PCR (Q-PCR) analysis in order to validate the changes detected by the microarray experiments. We focused on Pax6, expected to be downregulated both in anterior and posterior tissues of Raldh2−/− embryos, Id4, Six2 and Stat3 as examples of genes identified by microarray analysis as being downregulated anteriorly, and Lhx2, Crabp2, Bhlhe40 and Unc5a, that were scored as being downregulated in both regions of the mutant embryos. We also sought to confirm the upregulation of certain genes identified by microarray analysis, mainly for Bhlhe40 (Strat15) which, as discussed above, was not expected to be upregulated in the RA-deficient embryos. Creb5 (encoding a cAMP-responsive element binding protein) was chosen as an example of a gene with different behavior in anterior (slightly downregulated) versus posterior (upregulated) tissues.

Overall, the Q-PCR experiments showed a clear correlation with the changes observed by microarray analysis (Fig. 4). Of note, the Id4 and Six2 genes, that were scored as downregulated anteriorly in the microarray experiments, were also shown to be downregulated in posterior tissues by Q-PCR analysis, although with a lesser fold change than in anterior tissues. This could be explained by the higher sensitivity of the Q-PCR technique. The same observation was made for Stat4, which was downregulated posteriorly – to a lesser extent than in anterior tissues – according to Q-PCR analysis, whereas microarray analysis only revealed an anterior downregulation. In agreement with the microarray data, the Bhlhe40 gene was markedly upregulated, both in anterior and posterior tissues, according to Q-PCR results. Whereas Q-PCR showed a downregulation of Creb5 in anterior tissues of Raldh2−/− embryos, there was no significant change in transcript levels in posterior tissues.

Thus, for the selected genes the Q-PCR analysis tended to validate the microarray data. In particular, all genes that were scored as downregulated in the microarray analysis were confirmed by Q-PCR, the latter method often revealing a downregulation in the two types of tissue samples – while microarrays sometimes showed a change only in one type of samples. This points to some limitations in the sensitivity of DNA microarrays, implying that some genes identified as mildly downregulated could in fact be more severely affected in the mutant embryos.

We further attempted to document changes in expression by another technique, in situ hybridization (ISH). These experiments confirmed alterations in transcript levels and/or tissue distribution for some of the genes studied. Consistent with previous reports [29,34], Pax6 was downregulated in the caudal neural plate, as well as in more anterior regions of the developing neuroepithelium, including the forebrain, in early somite stage (E8.5) Raldh2−/− embryos (Fig. 5A). Lhx2 was downregulated in the forebrain of Raldh2−/− embryos, and was undetectable in posterior tissues expressing the gene in control embryos (Fig. 5B). An overall decrease of the Six2 ISH signal was observed in...
**Raldh2** 

2/2 embryos (Fig. 5C). Crabp2 was downregulated along the developing neural tube and caudal neural plate in early somite stage mutants (Fig. 5D, upper panels), and at later stages was also undetectable in neural crest-derived branchial arches cell populations (Fig. 5D, lower panels). Remarkably, **Bhlhe40** upregulation was also documented in **Raldh2**/2 embryos by ISH. At early somite stages, higher expression was detected in the hindbrain rhombomeres and extended ectopically along the neural tube (Fig. 5E, upper panels), whereas at later stages an abnormally high expression was observed in most regions of the embryo (Fig. 5E, lower panels). ISH experiments performed with **Id4**, **Stat4**, **Ikzf1** or **Unc5a** probes did not yield any consistent signal, either in wild-type or in **Raldh2**/2 embryos.

In summary, the changes detected by microarray analysis were for the most part confirmed by Q-PCR for selected genes, and could be documented at the tissue level for some of the genes by ISH. The latter technique has a limited sensitivity, as mRNA detection does not use an amplification step like in RT-PCR, and its readout is qualitative rather than quantitative. It is therefore not unexpected that some gene expression changes documented by transcriptome analysis and Q-PCR may not be visualized through ISH.

**Bioinformatic analysis of the transcriptomic data**

To get further insight into the cellular functions and classes of molecules that may be affected by RA deficiency, we performed an overall analysis of the transcriptome alterations using the Ingenuity Pathway Analysis (IPA) software. Figure 6 gives an overview of the main biological functions (with the more significant and therefore low p-values according to Ingenuity analysis) associated with the transcriptome of **Raldh2**/2 embryos. These functions are clearly relevant as they relate to general developmental processes (Cellular Development, Embryonic Development, Organ Development), or more specific phenomena for which retinoid signaling has already been implicated (Cardiovascular System Development, Skeletal and Muscular System Development for the POST microarray experiment). Inclusion of more basic biological functions (Cell-to-Cell Signaling and Interaction, Cellular Movement, Cellular Growth and Proliferation), as well as disease-related categories (Cancer, Neurological Disease, Genetic Disorder, Reproductive System Disease) are also not surprising considering the known effects of retinoids on cell growth and their use in clinical trials for chemotherapy or prevention of certain cancers [66]. These observations prompted us to further analyze the types

**Figure 7. Main classes of molecules downregulated in Raldh2−/− embryonic tissues, according to Ingenuity pathway analysis of the transcriptomic data.** Pie charts illustrate the distribution of gene products according to subcellular location (A) or protein function (B), when considering the overall set of affected genes (fold change ≥1.2, filtered for FDR <10% and a p-value ≤0.03 for POST and ≤0.05 for ANT). doi:10.1371/journal.pone.0062274.g007
of molecules and the molecular pathways affected by embryonic RA deficiency.

When the transcriptome data were sorted according to the subcellular localization of the corresponding proteins, there was an over-representation of nuclear proteins (mainly at the expense of cytoplasmic proteins) among genes affected in the POST microarray experiment (Fig. 7A). All major classes of molecules were represented among the sets of affected genes, both for the ANT and POST experiments (Fig. 7B). Notably, there was an over-representation of transcriptional regulators among affected genes in the POST experiment, whereas translational regulators were under-represented when compared to the ANT data set. A list of genes/proteins involved in transcriptional regulation and identified as down-regulated in the ANT and POST microarray experiments is provided as supplementary online information (Table S2).

Molecular pathways affected in Raldh2<sup>−/−</sup> embryos

Alteration of retinoic acid signaling has already been shown to interfere with several signaling pathways within the embryo (refs. [22,67,68] for reviews). Often the changes were described as changes in expression levels of ligands, such as FGFs [8,9], Wnts [69] or TGFβ [70]. In the case of the Sonic hedgehog (Shh) pathway, no change in ligand expression was observed in head or posterior embryonic tissue samples (fold change ±1.2, filtered for FDR <10%) is shown. The total number of genes comprising each pathway (middle line, gray shaded), and the percentages of genes misregulated in each experiment, are also given. doi:10.1371/journal.pone.0062274.g008

Figure 8. Summary diagram of the major molecular pathways emerging from Ingenuity analysis of the Raldh2<sup>−/−</sup> transcriptome. The molecular pathways are listed by decreasing order of statistical significance as they appear through analysis of the ANT (left-side list) and POST (right-side list) data sets. The most significant pathways identified for each data set are highlighted in gray. Additional pathways relevant for developmental processes are also listed. A graphic representation of the numbers of genes downregulated (green) or upregulated (red) in anterior (upper bars) or posterior (lower bars) Raldh2<sup>−/−</sup> embryonic tissue samples (fold change ±1.2, filtered for FDR <10%) is shown. The total number of genes comprising each pathway (middle line, gray shaded), and the percentages of genes misregulated in each experiment, are also given. doi:10.1371/journal.pone.0062274.g008

The molecular pathways that may be affected under conditions of embryonic RA deficiency. A graphic summary of the main pathways identified by Ingenuity analysis is provided in Figure 8. The molecular pathways are listed by decreasing order of statistical significance as they appear through analysis of the ANT (left-side list) and POST (right-side list) data sets. The six “top” (most significant) pathways for each data set are highlighted in gray, and additional pathways relevant for development are also illustrated. The numbers of genes downregulated (green bars) or upregulated (red bars) are illustrated (upper bars: ANT transcriptomic data; lower bars: POST transcriptomic data).

As observed through the analysis of biological functions, several “top” pathways emerging from the ANT microarray data are related to nervous system development or function (axonal guidance, Reelin signaling, Notch signaling, synaptic long term potentiation, dopamine receptor signaling). Although some of these phenomena take place at later developmental stages than those analyzed, their presence in the top list is easily explained as they contain genes also acting during early embryogenesis. The main molecular pathways identified by the analysis of POST microarray data are more varied, encompassing categories with large numbers of molecules (pyrimidine metabolism, purine metabolism, protein ubiquitination), and more specific pathways with a highly significant p-value, but with much fewer misregulated genes. For instance, the “role of Oct4 in pluripotency” pathway appeared among the top pathways with only 5 downregulated genes (Foxa1, Foxd3, Hoxb1, prohibitin Phb, and Polycystic ring finger Prg6). The “calcium signalling” pathway was characterized by the presence of 24 upregulated genes (vs. 1 downregulated genes, including those encoding alpha 1 (skeletal muscle) actin Acta1, cardiac myosin light chain 3 Myl3, and heavy chain 7 Myh7, calcium/calmodulin-dependent protein kinase IG CamkIg, Creb5, Nfatc4, Nfat3, and the histone deacetylase genes Hdac5 and Hdac8. Inclusion of the “one carbon (folate) metabolism” pathway among the most significantly affected ones may reveal a novel retinoid function, so far poorly studied, although some evidence for a regulation of this pathway by retinoids emerges from metabolomic studies in embryonic stem cells (L.J. Gudas, personal communication). The downregulated...
Table 2. Genes acting in selected signaling pathways and misregulated in Raldh2−/− tissue samples (ANT: anterior samples; POST: posterior samples).

| PATHWAY | ANT Upregulated | ANT Downregulated | POST Upregulated | POST Downregulated |
|---------|----------------|------------------|-----------------|---------------------|
| FGF     | Fgf10, Fgf11, Fgf16 | Fgf13, Fgf17 | Fgf10, Fgf11 |                     |
| Receptors | Fgf2 |                     |                 |                     |
| Effectors | Akt3, Gab1, Map2k1, Map2k6, | Hras, Mapk11, Pik3cb2, Pik3cb, | Akt3, Frs2, Map3k1, |                     |
|         | Map3k1, Map3k3, Met, Plcg1, Rac3 | Pik3r3 | Mapk13, Hras | Creb5 |
| BMP     | Bmp2, Bmp4 | Bmp2, Bmp4, Bmp5 |                     |                     |
| Receptors | Bmp1rb |                     | Bmp2 | Bmp1rb |
| Effectors/Modulators | Id1, Fst, Fstl3, Gpc3, Grem1, Htra1, Map2k1, Map3k, Mapk13, Msx2, Pitx1, Pitx2, Rras, Smad7 | Ecsit, Hras, Mapk10, Mapk11, Mras, Ptk4, Ptk5, Prkar1b, Rras, Shh | Mapk10, Msx2, Ptkx, Pitx2, Prkar1b, Rras, Smad7 | Hras, Mapk13 |
| TGFβ    | Inha, Tgfβ1, Tgfβ2, | Inha, Tgfβ1, Tgfβ2, |                     |                     |
| Receptors | AcvZa, Ptgfr, Tgfbr2, Tgfbr3 | Tgfbr2, Tgfbr3 |                     |                     |
| Effectors/Modulators | Ltbp1, Map2k1, Map4k1, Map3k, Rras, Smad6, Smad7, Tgfβ111, Tgfβ1, Tgfβ1 | Foxh1, Hras, Mras, Stat1 | Eng, Ltbp1, Rasa, Serpine1, Smad7, Tgfβ1, Tgfβ1, Tgfβ1, Thbs1 | Hras, Hraf, Tgfbrap1 |
| Hedgehog | Ihh |                     |                     |                     |
| Receptors | Ptc1 |                     | Ptc2 |                     |
| Effectors | Dyrk1b, Prkar1b, Hhip | Prkag2 | Prkar1b, Dyrk1b | Gli1 |
| IGF     | Ctfg, Cyr61 |                     |                     |                     |
| Receptors | Igif1 |                     | Igif1r |                     |
| Effectors/Modulators | Akt3, Igfbp2, Igfbp3, Igfbp4, Igfbp5, Iris1, Foxo3, Grb10, Map2k1, Map3k, Prkar1b, Rac3, Rras, Shc1 | Hras, Pik3c28, Pik3cb, Pik3r3, Prkag1, Prkag2 | Akt3, Foxo3, Igfbp2, Igfbp3, Igfbp5, Prkar1b, Rras, Shc1 | Hras |
| VEGF    | Fgf, Vegf, Vegfb, Vegfc | Pgf, Vegf, Vegfc |                     |                     |
| Receptors | Fli1, Fli4 |                     | Fli1 |                     |
| Effectors/Modulators | Actc1, Actg2, Actn3, Akt3, Foxo3, Map2k1, Map3k, Plcg1, Rac3, Rras, Shc1 | Elfay, Elf2b1, Elf2b4, Hif1a, Mras, Pik3cb, Pik3cb, Pik3r3 | Acta1, Actn2, Akt3, Foxo3, Kdr, Rras, Shc1 | Elf2b1, Elf2b4, Elf2s2, Hras |
| Wnt     | Wnt1, Wnt7a, Wnt7b | Wnt2, Wnt4, Wnt5a, Wnt7a |                     |                     |
| Receptors | Acv2a, Cdg2, Fzd5, Fzd7 | Cdg4, Fzrb, Ldr | Fzd4, |                     |
| Effectors/Modulators | Aes, Akt3, Cdh1, Cdh5, Dact1, Dkk1, Dkk3, Sostdc1, Calcocoi, Hbp1, Kremen1, Lrp1, Lrp5, Mif, Porcn, Ppp2r5c, Ppp2r3a, Ppp2r5c, Prkdc, Rac3, Ror2 | Apc2, Celsr2, Ndp, Ppp2r1b, Ppp2r2c, | Akt3, Calcocoi, Cdh1, Dakt, Dkk1, Dkk2, Gnao1, Hbp1, Lrp1, Mif, Ppp2r3a, Rspo1, Rspo3, Sostdc1, Tgf2, Tgf3, Wisp1 | Hnf1a, Lect2, Ndp, Ppp2r1b, Sfrp5 |
| Retinoic acid | Synthesis enzymes | Rdh10 |                     | Adh1 |
| Metabolism enzymes | Cyp26c1 |                     | Cyp26b1 |                     |
| Receptors | Rara, Rarg |                     | Rarg | Rarb |
| Effectors/Modulators | Cited2, Ncoa1, Nrf2f | Ncoa1, Nrc1, Rara2, Smarca2, Vegfa | Crabp2, Gdap1, Gdap2 | Crabp2, Raet1d1, Rbp1, Rbp4, Stra6, Trim12, Trt |

Key: **Bold** indicates a fold change exceeding ±1.5. 
*Italicics* indicate a p-value >0.05 (ANT) or >0.03 (POST). 

doi:10.1371/journal.pone.0062274.t002

Genes include Dhhf (dihydrofolate reductase) and Mthf1 (methylene tetrahydrofolate dehydrogenase). Furthermore, the folate receptor-coding gene Folr1 was markedly downregulated (−2.35), although with a p-value >0.05. Another potentially interesting pathway is that relating to CHK proteins in cell cycle control, which contained 7 downregulated genes, including Cdk23a, genes coding for E2F transcription factors 3 and 4 (E2f3, E2f4), Replication factor C3 (Rfc3), and Replication protein A1 (Rpa1).

Many of the signaling pathways acting during early embryogenesis appeared with a high significance value in the bioinformatic analysis of the transcriptomic data. This supports the idea of complex RA-dependent regulatory events, and “cross-talks” between the retinoid and other embryonic signaling pathways.
Interestingly, among the top (most significant) pathways listed for the ANT transcriptomic data, two were not the major candidates that would be predicted from the literature: VEGF signaling and ILK signaling. Three Vegf genes (Vegfa, Vegfb, Vegfc), as well as the Platelet-derived-growth factor C (Pdgfc) and Placental growth factor (Pgf) genes, appeared among the genes upregulated in both pathways (see Table 2 for a full list of misregulated genes in the VEGF signaling pathway). CDK5 and JAK/Stat signaling were also among the highest significant pathways in ANT tissues. Most affected genes in this category also appear as effectors in other pathways (see Table S2 for details). It is noteworthy to mention Stat4 as markedly downregulated (-3.07), Stat1 more mildly downregulated (-1.24), and Stat2, Stat5b and Stat6 upregulated in ANT tissues of mutants.

Table 2 gives details on genes misregulated belonging to several major embryonic signaling pathways (FGF, BMP, TGFβ, Hedgehog, IGF, Wnt/beta-catenin, and the retinoid pathway itself). In most cases the affected genes encoded for a subset of the ligands, more exceptionally a receptor (Fgfr2 upregulated in ANT tissues, Bmpr1b and Bmpr2, respectively downregulated and upregulated in POST tissues), and various effectors of the pathway (including Mapk genes acting in several pathways). While several Fgf genes were upregulated (Table 2), Fgf8, previously shown to be upregulated in RA-deficient embryos [2,9,10] was not picked up by the microarray analysis, an explanation being that the shift in the Fgf8 mRNA gradient [71] only occurs in a small region of the Raldh2−/− embryos (the anteriormost presomitic mesoderm). Upregulation of several Bmp genes – the highest for Bmp4 in ANT tissues, is also noteworthy. Within the TGFβ pathway there were a majority of upregulated genes, both in ANT and POST tissues. This is consistent with a previous study which identified the TGFβ pathway as being abnormally upregulated and involved in the phenotypic abnormalities observed during early embryonic lung development [70]. The Hedgehog signaling pathway exhibited the smallest numbers of misregulated genes (6 in ANT tissues, 4 in POST tissues), reflecting the small number of genes (28) included in this category, and included Gli1 (previously identified as downregulated in Raldh2−/− embryos: ref. [6]) and Indian hedgehog (Ihh, downregulated in POST tissues).

In summary, this bioinformatic analysis confirmed that retinoic acid deficiency affects the regulation of genes involved in a wide range of molecular pathways. Remarkably, the main developmental signaling pathways previously implicated as being retinoid-dependent (FGF, TGFβ/BMP, Hedgehog, Wnt; refs. [22,67,68] for reviews) do not appear as the top pathways in terms of statistical significance, hence the analysis summarized in Fig. 8.
provide new information on metabolic processes or signaling pathways that may deserve further investigation.

**Cross-comparison of in vivo (embryonic) data with results from genomic and transcriptomic screens in embryonic stem cells**

Several recent studies have used differentiating murine ES cells as a model for analyzing transcriptomic changes occurring after RA treatment [72,73,74], and/or for analysis of genomic loci bound by RARs, using large-scale chromatin immunoprecipitation (ChIP-ChIP or ChIP-seq) techniques [72,75,76]. We decided to perform a cross comparison of such studies, to identify which, among the genes downregulated in Raldh2−/− embryos, may also have been identified as being RA-inducible, or RAR-bound according to ChIP data, in differentiating ES cells. This comparison was performed with the data of Moutier et al., who reported the most extensive set of RAR-bound loci, identifying 6628 RAR occupied loci in ES cells grown for 4 days as embryoid bodies (EBs) and treated for 2 h with RA [72]. To narrow down the cross-comparison, their genomic list was restricted to loci in which the RAR-occupied sites were located within ±5 kilobases (Kb) from the transcription start site. These authors also performed a comparative transcriptomic (RNA-seq) analysis of EBs grown for 4 days and treated (or not treated) with RA, identifying 824 RA-induced transcripts [72]. We further used the data of Kim et al., who identified by Affymetrix array analysis 1202 gene transcripts upregulated in ES cells driven to a neural phenotype after exposure to RA from day 4 to 8 in a 12-day suspension culture [73]. Unekama et al. also reported transcriptional signatures of EBs driven to neural differentiation, and here we used their data set of 147 genes induced at day 8 of differentiation, after RA treatment from day 2 to 6 [74].

Figure 9 provides an overview of this comparative analysis in the form of Venn diagrams. An immediate major conclusion is that there are very limited overlaps between the sets of genes identified in this study, and those described as RA-inducible or RAR-bound in ES cells. Indeed there were only 31 (~2.2%) of the genes downregulated in anterior tissues, and 15 (~1.8%) of those downregulated in posterior tissues of Raldh2−/− embryos, that were tagged as RAR-bound by Moutier et al. [72]. Also, only 4% of the “anteriorly-affected” genes, and 5.5% of the “posteriorly-affected” genes appeared in the RNA-seq data of Moutier et al., i.e. behave as early RA-inducible genes in EBs. Additional genes were found by comparing our data to those of Kim et al. [73] and Akamuma et al. [74], which correspond to transcriptional signatures at a later stage of neural differentiation (2–4 days after the RA treatment). The complete gene lists are available as online supporting information (Table S3).

Surprisingly, only a handful (2 and 3 genes, respectively) were common to the three data sets, i.e. were RA-inducible and RAR-bound in ES cells, and affected anteriorly or posteriorly in Raldh2−/− embryos (Fig. 9). Whereas the anteriorly affected genes did not obviously appear as “developmental” genes (Atph1b1, ATPase, Na+/K transporter β1 polypeptide; Ascl4, acyl-coA synthetase family member 4; the retinoldehyde dehydrogenase/reductase gene Dhrs3) [39] was one such gene affected posteriorly. It may indeed be a key gene for feedback mechanisms regulating retinoid levels, both in the embryo and in cell systems. The few other genes common between our lists and the RAR-bound loci included several interesting candidates, namely the gene encoding fatty acid-binding protein (Fahp) 5 – identified in other cell systems as a regulator of the anti-apoptotic and proliferative response to RA [77], or genes encoding the BMP receptor Bmpq1b, the cell cycle regulator Ccarn1, and zinc finger (Znfx1, Zap420) or ring finger (Raf144b) proteins. Examples of genes downregulated in Raldh2−/− embryos, and reported as early RA-responsive in ES cells [72], are given in Fig. 9. While these include several known RA-target genes (described above), they highlight additional genes whose misregulation may be relevant for the RA-deficiency phenotypes.

**Analysis of cis-regulatory elements in downregulated genes**

Lastly, we used another bioinformatic approach to assess whether the genes defined here as being RA-dependent in the early embryo, may share other common regulatory features. We used the oPOSSUM online resource and database (refs. [78,79]; http://opossum.cisreg.ca/oPOSSUM3/) to perform this analysis. The oPOSSUM single site analysis (SSA) tool allows to identify over-represented transcription factor binding sites (TFBS) in specific sets of genes, giving scores to assess over-representation either at the gene level, or at the TFBS occurrence level. We interrogated the oPOSSUM database for SSA analysis on two sets of genes, corresponding to the most severely downregulated genes (FC < -1.5) in anterior or posterior Raldh2−/− tissues (146 and 63 genes, respectively). A summary of the retrieved data is provided in Table 3. Interestingly, the first ranked transcription factor (highest Z-score) for posteriorly downregulated genes is the RAR:RXR DR5 dimer. While this result further tends to validate our transcriptomic data, the corresponding gene hits were only 8, which included Dhrs3, Mox1, Metn, Rath, and Stat6. This finding would suggest that only a small fraction of the genes identified as downregulated in our study, are actually direct targets of RAR:RXR regulation.

Among other transcription factors with overrepresented binding sites are several forkhead factors (Foxa1, Foxa2 posteriorly, Foxa1, Foxd1, Foxd3, Foxo3 anteriorly). Binding sites for high mobility group (SRY-related or Sox) factors were also amongst the most overrepresented. Unlike RAR:RXR binding sites, these yielded very high numbers of hits, both in terms of gene numbers (in all cases >50% of the genes analyzed) and of TFBS identified (up to 782 per gene; see Table 3). This abundance probably reflects an overinflation of putative TFBS, often encountered with bioinformatic search engines, when the TFBS consensus is defined as a relatively short and degenerate sequence. The same conclusion was drawn for most of the homeodomain proteins identified in this search (e.g. Nobox, NKx2.5, Pbx1 or Hoxa5, all of which appeared with high Z-scores for both the anterior and posterior gene sets, and yield up to >1000 TFBS hits). This clearly points to a limitation of the search engine. However, and despite this overinflation, we would conclude that TFBS bioinformatic analysis will help selecting subsets of Raldh2−/− misregulated genes that may be regulated by specific homeodomain factors. A possible, combinatorial regulation by forkhead (Fox) and Sox proteins will also deserve further investigation.

On the other hand, the TFBS analysis provided several clues on regulation by specific transcription factors, with emerged with high Z-scores on much smaller numbers of putative targets. Noteworthy was the high Z-score of Pax5 and Pax6 for posterior genes (8 and 5 gene hits, respectively; Gad2, Dlx1, Rarb and Stat5a appearing as putative Pax6 targets). Regulation by Pbx1, a cofactor for Hox genes, was also highlighted for 15 posterior genes (Table 3). These included already documented Hox/Pbx targets such as Hoxa1 [80] and Rarb [81], and pointed to Ccarn1, Lgr5, Nep, NKx2-9 and 3-1 or Rfx6 as additional putative targets. A set of 8 genes was defined as putative Brachury (T) targets; these included Lbx1, Pnn2, Pdn13, NKx2-9 and Stat6. Involvement of STAT1 and/or STAT3 as possible regulators of several posterior genes was highlighted.
These included Ccne1, Crabp2, Dbx1, Dhrs3, Rarb and Stra6. Finally, three nuclear receptors were identified through this analysis: NR2F1 (COUP-TF I) as a putative regulator of 17 posterior genes, the PPARγ:RXRα dimer, with 22 putative targets among posterior genes, and NR3C1 (glucocorticoid receptor) with 21 identified targets among anterior genes (details on target genes in Table 3).

Altogether, this TFBS database search highlighted the combinatorial regulation of embryonic RA-dependent genes by several families of transcriptional regulators, while providing strong candidates for the regulation of specific subsets of genes. It also pointed out to four genes of the RA pathway: Dhrs3, Crabp2, Rarb, and Stra6, as key targets for such combinatorial regulation.

**Table 3.** Bioinformatic analysis of putative transcription factor binding sites (oPOSSUM3 single site analysis method) in genes showing the highest downregulation in posterior or anterior tissues of Raldh2−/− embryos.

| Transcription factor (a) | Family | Gene hits | TFBS hits | Z-score | Transcription factor (a) | Family | Gene hits | TFBS hits | Z-score |
|-------------------------|--------|-----------|-----------|---------|-------------------------|--------|-----------|-----------|---------|
| RXR:RAR (DR5) (b)       | Nuclear receptor | 8 | 11 | 11.40 | Box | Homeodomain | 107 | 679 | 21.90 |
| MZF1 5-13               | jlx; zinc finger | 43 | 272 | 10.27 | Nkx2-5 | Homeodomain | 124 | 1556 | 19.22 |
| Nkx2-5                  | Homeodomain | 48 | 262 | 9.90 | Pdx1 | Homeodomain | 120 | 1115 | 17.73 |
| Foxa1                   | Forkhead | 40 | 222 | 8.14 | Hoxa5 | Homeodomain | 123 | 1574 | 15.15 |
| Foxa2                   | Forkhead | 40 | 152 | 7.84 | REST (c) | jlx; zinc finger | 6 | 6 | 14.66 |
| Pax5 (d)                | Paired domain | 8 | 8 | 7.78 | Prx2 | Homeodomain | 109 | 938 | 13.98 |
| SRY                     | HMG group | 48 | 328 | 6.87 | Sox9 | HMG group | 88 | 379 | 12.99 |
| Nr2e1                   | Homeodomain | 52 | 606 | 6.86 | Sr | HMG group | 112 | 782 | 10.28 |
| Hnf1b                   | Homeodomain | 17 | 27 | 6.58 | Sox17 | HMG group | 109 | 558 | 10.16 |
| ARID3A                  | Arid | 48 | 494 | 6.08 | NfyA | Nfy CCAAT binding | 51 | 91 | 10.02 |
| Nr2f1 (e) (COUP-TF I)   | Nuclear receptor | 17 | 23 | 5.74 | Sox5 | HMG group | 99 | 554 | 8.86 |
| Stat1 (f)               | Stat | 19 | 40 | 5.49 | Foxd1 | Forkhead | 93 | 496 | 7.78 |
| Hnf1α                   | Homeodomain | 14 | 16 | 5.36 | Nkx3-1 | Homeodomain | 103 | 530 | 6.63 |
| Pdx1                    | Homeodomain | 49 | 418 | 5.35 | Foxd3 | Forkhead | 85 | 340 | 5.86 |
| Pbx1 (g)                | Homeodomain | 15 | 25 | 5.21 | Pou5f1 (Oct4) (h) | Nuclear receptor | 26 | 40 | 5.81 |
| Pparγ:Rxrα              | Nuclear receptor | 22 | 40 | 3.96 | Lhx3 | Homeodomain | 41 | 88 | 5.56 |
| Hoxa5                   | Homeodomain | 52 | 607 | 3.70 | Foxa3 | Forkhead | 95 | 537 | 5.47 |
| T (Brachyury) (i)       | J-ribbon | 8 | 10 | 3.63 | Nnr3c1 (Gf) (j) | Nuclear receptor | 21 | 28 | 5.12 |
| Stat3                   | Stat | 32 | 82 | 3.48 | Hnf1b | Homeodomain | 33 | 54 | 4.70 |
| Pax6 (k)                | Paired domain | 5 | 6 | 3.26 | Foxa2 | Forkhead | 82 | 323 | 3.97 |

(a)59 genes analyzed from 63 genes downregulated with FC < −1.5.
(b)137 genes analyzed from 146 genes downregulated with FC < −1.5.
(c)The searched region encompassed 5 Kb upstream and 5 Kb downstream of the transcription start site. The top ten factors (highest Z-scores) appear in bold. Additional transcription factors (below) are the authors’ selection.
(d)Genes with conserved RAR/RXR (DR5) binding sites are: Ccn3, Dhrs3, Fap, Mefx1, Metnr, Rarb, Strad, 573056820Rik (NB: 573056820Rik is an antisense EST found between the Hoxa3 and Hoxa4 genes).
(e)Genes with conserved Pax5 binding sites are: Apba2, Crabp2, Lgr5, Mettl11, Nkx2-9, Pmm2, R hippy3, Rfx6.
(f)Genes with conserved Nkx2-1 binding sites are: Apba2, Crabp2, Dhrs3, Dusp9, Fap, Gchb, Hoxa1, Maob, Nkx2-9, Pdx, Prdx13, Rarb, Sfrp5, Stra6, Timm8a1, Tmem56, 1700011H4Rik.
(g)Genes with conserved TFBS analysis: genes downregulated in POSTERIOR tissues (a).

Conclusion

This study provides an overview of the genes misregulated in early mouse embryos unable to synthesize retinoic acid (RA) from its precursor retinaldehyde, due to targeted gene disruption of a critical biosynthesis enzyme. While highlighting several genes known to be affected by conditions of impaired RA signaling in mouse embryos and/or other animal models, it provides additional candidate genes and pathways that may be relevant to better understand the mechanisms underlying the complex
phenotypic consequences of embryonic RA deficiency. Bioinformatic analysis confirmed the impact of retinoid deficiency on several signaling pathways, with specific genes misregulated and belonging to the FOH, Notch, BMP, Hedgehog, VEGF, or Wnt pathways. This analysis pointed out to additional pathways or biological functions which, although not previously correlated with retinoid deficiency, appeared in the top (most significant) categories. This provides clues for further exploration of the effect of retinoid deficiency on metabolic processes such as purine and pyrimidine metabolism or one carbon metabolism, on signaling pathways such as the mTOR, ILK and IGF1 pathways, or on protein ubiquitination. Our data also suggest an upregulation of calcium signaling, specific to posterior (trunk) tissues of RA-deficient embryos.

This study adds to a framework of other high-throughput studies that reported RA-dependent transcriptional alterations [72,73,74,82], or characterization of RAR target gene loci [72,75,76,82], in tissue culture cells including embryonic stem cells. Remarkably, a cross comparison between several data sets revealed that only a small subset of genes downregulated in Raldh2−/− embryos were previously identified as RAR-bound, or RA-inducible, in embryonic stem cells. Future high-throughput genomic/transcriptomic studies, expected to be performed on other retinoid target tissues, should improve our knowledge on retinoid-dependent gene regulation, both during normal development and physiological processes, and with respect to various diseases including cancer.

Materials and Methods

Ethics statement, mouse lines and embryonic tissue collection

Animal experimentation protocols were reviewed and approved by the Direction Départementale des Services Vétérinaires (agreement # 67–189 dated March 30, 2009) and conformed to the NIH and European Union guidelines, provisions of the Guide for the Care and Use of Laboratory Animals, and the Animal Welfare Act. No invasive (pain-inducing) procedure was used for this study. The mice harbored the Raldh2 knockout allele described in ref. [26] and, for the X-gal assays, the RARE-hsp68-lacZ transgene described in ref. [32]. Pregnant female mice were euthanized after anesthesia with CO2, embryos were collected and transgene described in ref. [32]. Pregnant female mice were euthanized after anesthesia with CO2, embryos were collected and transgene described in ref. [32]. Pregnant female mice were euthanized after anesthesia with CO2, embryos were collected and transgene described in ref. [32]. Pregnant female mice were euthanized after anesthesia with CO2, embryos were collected and transgene described in ref. [32]. Pregnant female mice were euthanized after anesthesia with CO2, embryos were collected and transgene described in ref. [32]. Pregnant female mice were euthanized after anesthesia with CO2, embryos were collected and transgene described in ref. [32]. Pregnant female mice were euthanized after anesthesia with CO2, embryos were collected and transgene described in ref. [32].

In situ hybridization and X-gal staining

Whole-mount in situ hybridization with digoxigenin-labeled riboprobes was performed as described [83], using an Intavis InSituPro robot (for details, see http://www.empress.har.mrc.ac.uk/browser/, Gene Expression section). X-gal assays were performed as described [32].

Supporting Information

Figure S1 Hierarchical clustering of differentially expressed genes. The heatmap shows the gene expression profiles of microarray data sets (upregulated expression: red; downregulated expression: green). Importantly, the WT and KO samples segregate into fully distinct clusters according to this analysis. ANT: RNA from anterior tissues; POST: RNA from posterior tissues; WT: wild-type embryos; KO: Raldh2−/− embryos. (TIF)

Table S1 Genes exhibiting a “contradictory” behavior in the microarray experiments, i.e. downregulated in one type of tissue samples and upregulated in the other (±1.2 fold change) in Raldh2−/− embryos. (DOC)

Table S2 Genes involved in the process of transcription/transcriptional regulation and downregulated in Raldh2−/− embryos. Genes are listed separately for the ANT and POST microarray experiments. Genes with <-1.5 fold changes appear in bold. (PDF)

Table S3 Genes found in common in our transcriptomic study (genes downregulated in anterior or posterior tissues of Raldh2−/− embryos), and in studies which identified RA-inducible genes by RNA-seq (ref. [72]) or...
microarray (refs. [73,74]) analysis of differentiating ES cells. A cross-comparison with genes reported as RAR-bound in differentiating ES cells (ref. [72]) was also performed.

(PDF)

Table S4 Sequences of primers used in the quantitative RT-PCR experiments (F: forward primer; R: reverse primer).

(PDF)

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Acknowledgments

We thank Valérie Fraulob, Brigitte Schulhaub, and members of the IGMBMC microarray core facility, for their technical assistance. We are grateful to Drs. Irwin Davidson, James Huettner, and Hideki Sone, for kindly providing transcriptional/genomic data sets.

Author Contributions

Conceived and designed the experiments: MP KN PD. Performed the experiments: MP CS MR CT. Analyzed the data: MP DD PD. Wrote the paper: MP KN PD.
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