The Transcriptional Response to Oxidative Stress during Vertebrate Development: Effects of tert-Butylhydroquinone and 2,3,7,8-Tetrachlorodibenzop-\(\beta\)-Dioxin

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Abstract

Oxidative stress is an important mechanism of chemical toxicity, contributing to teratogenesis and to cardiovascular and neurodegenerative diseases. Developing animals may be especially sensitive to chemicals causing oxidative stress. The developmental expression and inducibility of anti-oxidant defenses through activation of NF-E2-related factor 2 (NRF2) is affected by developmental stage, and may vary among species. In zebrafish, there is substantial overlap between genes induced in developing zebrafish and a set of marker genes induced by oxidative stress in mammals. Genes induced by tert-Butylhydroquinone (tBHQ) as early as 1 dpf were significantly different among the DMSO-, tBHQ-, and TCDD-treated eleutheroembryos at 4 dpf. There was substantial overlap between genes induced in developing zebrafish and a set of marker genes induced by oxidative stress in mammals. Genes induced by tBHQ in 4-dpf zebrafish included those involved in glutathione synthesis and utilization, signal transduction, and DNA damage/stress response. The strong induction of hsp70 determined by microarray was confirmed by qRT-PCR and by use of transgenic zebrafish expressing enhanced green fluorescent protein (EGFP) under control of the hsp70 promoter. Genes strongly down-regulated by tBHQ included mitfa, providing a molecular explanation for the loss of pigment in tBHQ-exposed embryos. These data show that zebrafish embryos are responsive to oxidative stress as early as 1 dpf, that responsiveness varies with development in a gene-specific manner, and that the oxidative stress response is substantially conserved in vertebrate animals.

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Introduction

Oxidative stress occurs when redox signaling and control are disrupted, either through generation of non-physiological levels of reactive oxygen species (ROS) or by alterations in the regulation of key thiol/disulfide couples [1]. Oxidative stress has been cited as a causative or contributing factor in a variety of human conditions linked to environmental exposures, ranging from chemical teratogenesis to cardiovascular and neurodegenerative diseases [2–6]. Redox-mediated signaling is thought to be important for cellular differentiation during embryonic development [7,8] and developing embryonic stages may be especially sensitive to the disrupted redox and sulfhydryl balance that characterizes oxidative stress [9–12]. Oxidative damage has been implicated in the
mechanism of action of several developmental toxicants, including known human teratogens (e.g. thalidomide, phenytoin, ethanol), environmental contaminants (e.g. 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD), benz[a]pyrene), and nanomaterials [9,13,14], as well as in the etiology of congenital malformations associated with diabetic embryopathy [15,16].

The constitutive (basal) expression and inducibility of anti-oxidant defenses are known to affect the susceptibility of adult tissues and cells to effects of oxidative stress [17,18], and are likely to be important determinants of susceptibility at early life stages as well [19,20]. In adult animals, oxidant and pro-oxidant chemicals elicit an oxidative stress response (OSR), which involves the increased expression of genes whose products act to mitigate the oxidant challenge. Oxidants, electrophiles, sulfhydryl-reactive agents, and some phenolic anti-oxidants initiate this response by activating NF-E2-related factor 2 (NRF2) [NFE2L2] and related cap’n’collar (CNC)-basic-leucine zipper (bZIP) family proteins. (For nomenclature conventions, please see footnote 3 of reference [21].) NRF2 is normally found in the cytoplasm, where an interaction with Kelch-like-ECH-associated protein (KEAP1) targets it for rapid proteasomal degradation [22]. Oxidative stress disrupts the interaction between NRF2 and KEAP1, after which NRF2 enters the nucleus and forms a heterodimer with one of several small MaF proteins; NRF2-MaF dimers bind to anti-oxidant response elements (ARE) and activate transcription of genes such as glutathione S-transferases (GST), NAD(P)H-quinone oxidoreductase (NQO1), glutamate-cysteine ligase (catalytic subunit; GCLC), heme oxygenase (HMOX), and superoxide dismutase (SOD) [18,23,24].

Despite the demonstrated importance of the OSR in adults, the ability of vertebrate early life stages to respond to oxidative insult is not well understood. How does the sensitivity of developing vertebrates to oxidative stress vary with developmental stage? Are the patterns of induced or repressed gene expression stage-specific? What transcription factors are involved in regulating the OSR in embryos?

To begin to address these questions and elucidate the fundamental mechanisms by which vertebrates respond to oxidative stress during development, we have initiated studies to identify the core set of genes and the transcription factors involved in the OSR in developing zebrafish. The zebrafish (Danio rerio) is an established model in developmental biology that has emerged also as a valuable in vivo system in which to examine mechanisms of toxicity in developing animals and to screen chemicals for regulation in zebrafish remain unanswered. First, the ontogeny of the OSR in zebrafish embryonic and larval stages is not well understood. Kobayashi et al. [33,39] reported that tert-butylhydroquinone (tBHQ) can induce an OSR in zebrafish larvae (4 to 7-dpf), but that embryos at 6–24 hpf were incapable of mounting an acute OSR (as measured by induction of gsp1). Timme-Laragy et al. [30] observed an OSR (induction of gsp1, gpx1, and gclc) in zebrafish embryos exposed to tert-butylhydroquinone or 2,3,7,8-tetrachlorodibenzo-p-dioxin for 24 hr beginning at 24-hpf. These reports suggest that there are stage- or chemical-specific differences in responsiveness to oxidative stress, highlighting the need for a more systematic assessment of the basal (constitutive) and oxidant-inducible expression of OSR genes during development. Second, the set of oxidant-responsive genes in adult mammals or mammalian cell culture has been studied extensively (e.g. [47,48]), but the overlap between the OSR in zebrafish and mammals, and the identity of genes that respond to oxidative stress in developing vertebrates, are not yet well understood.

Here we report the results of studies using expression profiling to assess the nature of the response of developing zebrafish to chemically mediated oxidative stress. We conducted experiments in which gene expression was measured in zebrafish exposed to tBHQ at various times during development, including embryos (1, 2, and 3-4 dpf), clethroembryos (4 and 5-6 dpf), and larvae (6-7 dpf) (developmental phases according to [28,49]; Fig. 1). tBHQ is widely used as a prototypical mono-functional inducer of the OSR [23]. For comparison and to differentiate between NRF- and AHR-mediated responses, we also exposed developing zebrafish to TCDD, which is well known to cause altered gene expression through activation of AHRs in zebrafish embryos [50,51], and has been suggested to cause embryotoxicity at least in part by causing oxidative stress [52]. The results identify both similarities and differences between the OSR in developing zebrafish as compared with that elicited by tBHQ in previous studies in adult mammals or mammalian cells. In addition, we identify a link between a specific change in gene expression (repression of mitfa) and a phenotypic response to tBHQ [loss of pigmentation] and we describe the potential use of an existing transgenic line of zebrafish [53] to further investigate the temporal and spatial regulation of the anti-oxidant response in developing vertebrates.

Results
Expression and induction of oxidative stress response genes in zebrafish embryos
To assess the ability of embryos to mount a response to oxidative stress, we performed an experiment in which separate groups of embryos, clethroembryos, or larvae at 1, 2, 3, 4, 5, and 6-dpf were exposed for 6 hr to DMSO, tBHQ (10 μM), or TCDD (2 nM) and gene expression was measured by qRT-PCR and microarray. The goal of this experiment was to assess the acute response (6 hr) to chemical treatment occurring at different stages of development (Fig. 1), rather than secondary changes that might occur after a longer exposure time.

Targeted analysis of gene expression by qRT-PCR showed that tBHQ caused strong (4- to 8-fold) induction of gsp1 in 1- and 2-dpf embryos, but not at later stages (Fig. 2). tBHQ also induced gclc (3- to 4-fold; 1- and 4-dpf) and nrf2a (1- and 2-dpf), but not sod1, nqo1, or cyplα. TCDD induced cyplα at all time points (100- to 600-fold) and nrf2a at 2- and 3-dpf (2- to 3-fold), but did not significantly alter the expression of the other genes examined, under these exposure conditions. The induction of nrf2a expression by TCDD is consistent with recent studies showing regulation of NRF2 expression by the AHR in mammals [54,55]. Overall, the data demonstrated that some classical OSR genes (gsp1, gclc) were responsive to tBHQ-induced oxidative stress in embryos as early as 1- and 2-dpf, while others (sod1 and nqo1)
were not inducible by a 6-hr exposure to this concentration of tBHQ (10 μM) at any of the developmental stages examined here.

To more comprehensively assess the set of oxidant-responsive genes in zebrafish embryos, we examined a subset of the RNA samples by microarray using the Agilent 22k zebrafish array. We analyzed all of the 4-dpf samples, which included four biological replicates each for DMSO-, tBHQ- and TCDD-treated embryos. The 4-dpf time was chosen for gene expression profiling because Kobayashi et al. [33,39] had reported a robust induction of several oxidant-responsive genes by tBHQ in zebrafish larvae at this stage. Probes indicating significantly different relative transcript abundance among the DMSO, TCDD, and tBHQ treatments were determined by ANOVA, with the false discovery rate (FDR) set to 5%. Overall, 1477 probes exhibited significant differences in expression among the three exposure groups. For each of the probes we calculated fold-change in gene expression for TCDD- and tBHQ-treated embryos versus DMSO-treated embryos (Table S2).

FatiGO+ [56] was used to examine enrichment of Gene Ontology (GO) terms in the set of significant probes relative to the entire microarray probe set. GO terms enriched in the set of probes detecting transcripts that were up-regulated in response to tBHQ included “glutathione metabolic process”, “response to temperature stimulus”, “protein dimerization activity”, and several categories involving ester hydrolases and protein phosphatases (Table S3). Down-regulation of transcript abundance by tBHQ was associated with enrichment of GO terms for “negative regulation of cellular process”, “regulation of transcription”, “transcription”, and several categories of proteases (Table S3).

(The ability to perform GO analyses was limited by the incomplete annotation of probes on the Agilent array and by incomplete GO annotation of the zebrafish genome. Thus, when interpreting the FatiGO+ results it is important to consider the current status of zebrafish genome annotation. FatiGO+ relies on the assignment of GO terms within the Ensembl annotation of the zebrafish genome. However, many Agilent microarray probes...
have yet to be assigned to transcripts or genes at Ensembl and are thus not assigned GO terms. Others are assigned to transcripts or genes with preliminary annotation and GO term assignment only and are thus relatively uninformative, restricting Fatigo+ to detection of broad GO terms only (e.g. catabolic process, transcription, protein dimerization activity).

To prioritize the 1477 genes with significant changes for further study, we chose to focus only on those genes exhibiting ≥2-fold change in expression (up or down) for either TCDD or tBHQ, of which there were 345 (Table S4). This additional filter yielded 220 probes that were both significant in the statistical analysis and ≥2-fold up-regulated by tBHQ, and 109 that were significant and ≥2-fold down-regulated by tBHQ. For TCDD, 17 of the significant probes were ≥2-fold up-regulated and 8 were ≥2-fold down-regulated. There was very little overlap in probes responsive to tBHQ and TCDD (Fig. 3). At the ≥2-fold level, four probes (two hspl70 probes, dnah1, hspl90a2) were up-regulated by both tBHQ and TCDD and two probes were down-regulated by both compounds; one probe (foxp1b) was up-regulated in response to TCDD but down-regulated in response to tBHQ, while two probes were down-regulated in response to TCDD but up-regulated in response to tBHQ. After all annotation efforts, 14 of the 345 probes (4.1%) did not have a significant match to known genes or expressed sequence tags (ESTs).

Genes altered in response to tBHQ: Conserved vertebrate response to oxidative stress

One goal of this study was to compare the transcriptional response of developing zebrafish to a model oxidant (tBHQ) with the response that has been documented previously in adult mammals or mammalian cell lines. Any discrepancies might suggest either fish-mammal or embryo/larval-adult differences in the nature of the oxidative stress response. We therefore compared the set of genes induced by tBHQ in our experiment (Table S2) to a set of 18 “biological marker” genes that are induced by oxidative stress in mammals and mammalian cells through activation of the NRF2-ARE pathway [47]. Of the 18 markers, homologs of ten (56%) were up-regulated in 4-dpf zebrafish eleutheroembryos, five were unchanged, and three were not present on the array (Table 1). Several of the marker genes occur as duplicates in zebrafish; for four of these (hspl90, dnah1, thioredoxin, ferritin heavy polypeptide), only one of the paralogs was induced by tBHQ.

To further assess the nature of the eleutheroembryo response to tBHQ, we looked for altered expression of genes involved in the synthesis and utilization of glutathione (GSH) and other sulphhydryl-reactive anti-oxidants (thioredoxin, peroxiredoxin) and their regulation (Table 2). Among the induced genes, those associated with GSH synthesis included gcl, gcll, and glutathione synthase (gss). Induced genes associated with GSH utilization included GSH reductase (gr1), gamma-glutamyl transferase (ggt1a), GST omega, and a microsomal GST. Thioredoxin (txn1) and cystathionine beta-synthase (cbs), which are involved in cysteine disulfide reduction and cysteine synthesis via transsulfuration, respectively, also were induced by tBHQ.

There was strong induction of keap1a and several genes encoding small mal proteins (mafk, mafi, mafj), which are involved in the NRF2 signaling pathway. In contrast, four of the six predicted NRF-family transcription factors (nfe2, nrf1a, nrf1b, and nrf2a) were represented on the array, but none showed significantly altered expression in tBHQ-exposed eleutheroembryos.

Another set of genes of interest was those involved in general stress responses. The gene induced most highly by tBHQ (more than 50-fold) was hspl70, which was represented by several probes on the array (Table 3). A v-fos homolog was also strongly induced (47-fold); there were also substantial increases in expression of junb (3.7-fold) and other jun-related transcripts. Other induced genes included an hspl90 isoform (hspl90-alphain2) and hspl40/duaf1. The latter joined alf3 and gadl4/5 (several forms) as induced genes associated with a DNA-damage response.

cis-regulatory elements.

Among the genes significantly up-regulated by tBHQ exposure in the microarray experiment, some contained a possible DRE (e.g. early growth response 2a, arrestin domain containing 2, angiotensinogen, dual specificity phosphatase 5, phosphogluconate dehydrogenase isoform 1), an NF-kappaB motif (e.g. kelch-like ECH-associated protein 1a, growth arrest and DNA-damage-inducible, beta), SP1 motif (arrestin domain containing 2, pyruvate dehydrogenase kinase 2, dual specificity phosphatase 5, hypoxia induced gene 1, mmp13), REL motif (kelch-like ECH-associated protein 1a, growth arrest and DNA-damage-inducible beta, myocyte enhancer factor 2d), the MaB motif (e.g solute carrier family 25 member 43), and HIF1A::Arnt motif (e.g. heat shock cognate 70-like, solute carrier family 25 member 43).

For genes significantly down-regulated by tBHQ exposure, a number contained a SP1 motif (e.g. forhead box D1-like), growth arrest-specific 1a, red-sensitive opsin-1, microphthalmia-associated transcription factor a, insulin-like growth factor 2a precursor) and a MaB motif (rad and gem-related GTP-binding protein 1, aquaporin 3a, red-sensitive opsin-1).

Novel responses to tBHQ.

In addition to the stress-responsive genes and those involved in GSH homeostasis and other adaptive responses to oxidative stress, there were several notable changes in expression of other genes, including some with important roles during development (Table 3; Table S2). For example, sox9a, which has roles in development of chondrocytes and the retina [57,58], was induced 2.4-fold by tBHQ. The fibroblast growth factor inhibitor sprouty4 was induced 5-fold. Transcripts for the iron-regulatory protein hepcidin (hepcidin antimicrobial peptide; hamp1) were induced 8.3-fold. Prostaglandin endoperoxide synthase 2a (ptgs2a; also known as cox2) was induced almost 6-fold by tBHQ. Hypoxia-induced gene (big1) was induced 7-fold. Several members of the solute carrier family (slc25a25, slc16a9a, slc16a3, slc1a4, slc25a43, slc16a6b and slc13a2) were strongly induced, suggesting a general up-regulation of transport activity. Several dual-specificity phosphatases (dusp5, dusp4, dusp1) also were induced.

Although reports of the response to oxidative stress often emphasize the genes that are induced, oxidative stress also leads to decreases in the expression of some genes. In zebrafish eleutheroembryos exposed to tBHQ at 4 dpf, almost a third of the genes with significant and ≥2-fold changes were down-regulated, several strongly so (Table S2). A number of these suggest effects on the eye. For example, opsin 1 (opn1lw1), microphthalmia-associated transcription factor a (mitfa), and genes involved in retinoid homeostasis (lecinthin retinol acyltransferase a, retinol binding protein 4) were among those strongly suppressed by tBHQ exposure at 4 dpf. Several genes with known developmental roles also were repressed by tBHQ. Examples include foxp1b, frizzled homolog 2, kruppel-like factor 2a, distal-less homeobox gene 3b, lunatic fringe, noggin1, and fibroblast growth factor 8 (Table 4).

Genes altered in response to TCDD.

The number of genes affected by TCDD was small in comparison to the number regulated by tBHQ. Not surprisingly, cyp1a exhibited the greatest degree of induction (>70-fold), confirming the effectiveness of the exposure regimen. hspl70 also was induced by TCDD, but the
induction was modest (less than 4-fold) as compared to that caused by tBHQ (>50-fold). The forkhead box gene foxq1b was induced 3.5-fold by TCDD. Other TCDD-induced genes included ugt1b5 and ahr2 (Tables 2 and 3; see also Tables S2 and S4). For those genes responding to TCDD, we detected cis-regulatory elements in Ahr2 (ARE, Arnt::Ahr, Mofb, HIF1A::Arnt), CYP1A
Table 1. Zebrafish vs mammalian response to tBHQ.

| Marker gene in mammals | Zebrafish (co-)orthologs | # probes | tBHQ/DMSO |
|------------------------|--------------------------|----------|-----------|
| HSP70 1A, 1B, 6        | hsp70 (Chr.3)            | 2        | 52.3±13.8 |
|                        | hsp701 (Chr.8)           | 1        | 15.5±4.50 |
| HSP90 9B               | hspa9 (Chr.14)           | 2        | 1.19±0.22 |
| HSP90 1 alpha          | hsp90a (Chr.20)          | 3        | 1.18±0.32 |
| DnaJ (Hsp40) B1        | dnakb1a (Chr.3)          | 3        | 1.39±0.07 |
|                        | dnakb1b (Chr.1)          | 1        | 10.48±2.33|
| Heme oxygenase-1 (HMOX1)|                         | 0        | NA        |
| NADPH quinone oxidoreductase-1 (NQO1) | nqo1 | 1 | 0.66±0.08 |
| Glutamate-cysteine ligase, modifier subunit | gclm | 1 | 4.11±0.18 |
| Thioredoxin (TXN)      | txn1 (Chr.7)             | 2        | 5.47±0.92 |
|                        | txn2 (Chr.1)             | 2        | 0.83±0.21 |
| Thioredoxin reductase-1| txmrd1                   | 1        | 1.56±0.14 |
| Malic enzyme 1         |                          | 0        | NA        |
| Glutathione reductase  | glutathione reductase 1  | 2        | 3.79±0.68 |
| Ferritin, heavy polypeptide-1 | fth1 (Chr.7) | 5 | 0.98±0.04 |
| Ferritin-like (Chr.3)  | ftrl-like (Chr.3)        | 2        | 6.81±1.14 |
| Ferritin-like (Chr.25) | ftrl-like (Chr.25)       | 1        | 0.66±0.06 |
| Ferritin light polypeptide | ftrl                  | 3        | 0.89±0.10 |
| Carbonyl reductase-1   | cbr1                     | 2        | 1.09±0.27 |
| Phosphogluconate dehydrogenase | pgd | 2 | 2.46±0.40 |
| Sequestosome-1         | sqstm1                   | 1        | 7.57±1.56 |
| Ubiquitin thioesterase | usp4                     | 2        | 1.00±0.08 |
| Spermidine/spermine N1-acetyltransferase |             | 0        | NA        |

Set of 20 candidate genes based on the multiple data sets of genes responding to oxidative stress in mammalian cells, compiled by Johnson et al, as listed in Table 2 of Li et al Physiol Genomics 21:45-58, 2005. The 20 mammalian candidate genes have been collapsed into 18 sets based on orthologous or co-orthologous relationships with zebrafish genes. Of 18 genes, 10 have at least one co-ortholog induced by tBHQ in 4-dpf zebrafish (indicated in bold type), 5 are not induced, and 3 are not found on array.

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(ARE, Arn::Ahr, HIF1A::Arnt, SP1), and 5' nucleotidase, cytosolic II, like 1 (ARE, Arn::Ahr, SP1).

Analysis of selected genes by qRT-PCR. To confirm the changes in gene expression measured by microarray and to further explore the timing of those changes with respect to zebrafish development, we considered the six genes originally chosen for targeted analysis by qRT-PCR (gstp1, gclc, sod1, cyp1a, nqo1, nr1f2a; Fig. 2) and measured the expression of six additional genes, including both tBHQ up-regulated (hsp70, gadd45, atf3) and tBHQ down-regulated genes (mitfa, opn11u1, foxl1b) (Fig. 4). There was excellent concordance between microarray data and day 4 qRT-PCR data (Table S5). Comparing the direction and statistical significance of the responses to tBHQ and TCDD for the twelve genes examined (i.e. 24 comparisons), there was agreement in all but one case: the decreased expression of foxl1b in tBHQ-exposed eleutheroembryos was not reproduced by qRT-PCR. Notably, gstp1 showed a trend towards induction in both analyses (and was induced at other time points), but the high biological variation in the day 4 samples precluded statistical significance for both array data and qRT-PCR results. For all genes, the magnitude of change measured by qRT-PCR was as great or greater than that seen on microarray, in agreement with the well-known compression of fold-change values observed with array data [59]. Nevertheless, the confirmation of 23/24 microarray-detected changes by qRT-PCR supports the use of this platform for initial identification of tBHQ- and TCDD-induced changes in gene expression in developing zebrafish.

The qRT-PCR measurements also revealed varied developmental patterns of sensitivity to altered gene expression in response to tBHQ or TCDD during the first six days of development. Some genes were induced (hsp70, gadd45a, atf3, cyp1a) or repressed (mitfa) at all stages examined, whereas for other genes the altered expression occurred only at certain stages (gstp, gclc, nr1f2, opn11u1, foxl1b) or not at all (nqo1, sod1).

Loss of pigmentation in embryos exposed to tBHQ linked to altered expression of mitfa

A recent report [60] noted briefly that zebrafish embryos exposed to tBHQ exhibited reduced pigmentation, an observation that was confirmed in our studies. To explore the timing and persistence of the effect of tBHQ on pigmentation, we exposed embryos to tBHQ (5 or 10 μM) from 32–47 hpf and examined pigmentation at 52 hpf. In DMSO-exposed embryos, melanophores could be observed in their normal positions near the otic vesicle, in the dorsal and ventral stripes, and around the yolk sac (Fig. 5A). In embryos exposed to tBHQ, melanophores in the head and trunk were present but were small and hypopigmented; melanophores were not apparent on the yolk sac. In addition, tBHQ-exposed embryos displayed a reduction in pigmentation in the retina. By 120-hpf (5-dpf), 3 days after tBHQ exposure had
ended, there was partial recovery of pigmentation in the dorsal, lateral, and ventral stripes, in the yolk sac, and in the retina (Fig. 5B).

One of the genes most strongly repressed in tBHQ-treated eleutheroembryos was *mitfa*, one of two zebrafish co-orthologs of the mammalian gene microphthalmia-associated transcription factor (MITF) [61]. The *mitfa* gene is defective in zebrafish *nacre* mutants, which lack melanophores and thus do not synthesize melanin other than in the retina [62]. The reduced expression of *mitfa* suggests a molecular explanation for the reduced pigmentation in tBHQ-treated embryos.

### Response of *hsp70-EGFP* transgenic zebrafish to tBHQ

There was a strong induction of *hsp70* at 4 dpf in response to tBHQ exposure (Table 3, Fig. 4) and studies in human cells have shown that *hsp70* expression can be regulated by NRF2 [63]. Together, these results suggested that the *hsp70* gene might serve as a useful marker for induction of the oxidative stress response during development.

Kuwada and coworkers [53] developed a transgenic line of zebrafish containing the gene for enhanced green-fluorescent protein (EGFP) under control of the zebrafish *hsp70* promoter. In un stressed fish, EGFP is expressed only in the lens of the eye [64]. To assess the response of the *hsp70-egfp* fish embryos to oxidative stress, we exposed them to tBHQ and looked for induction of EGFP. At 24 hpf, tBHQ (10 μM) induced EGFP expression in a restricted pattern of cells including the eye and in specific cells in the trunk (Fig. 6A). When *hsp70-EGFP* eleutheroembryos were exposed to tBHQ (10 μM) for 4 hr beginning at 96-hpf and examined 4 hr later (104 hpf, similar to the sampling point for our microarray studies), tBHQ induced widespread EGFP expression throughout the fish (Fig. 6B), consistent with the high level of induction of *hsp70* measured by microarray and qRT-PCR.

### Discussion

The ability of embryos to protect themselves against oxidative damage is critical for maintaining developmental processes in the face of exposure to chemicals that are capable of disrupting redox balance and sulfhydryl metabolism. Despite the importance of such protective mechanisms, the ontogeny of constitutive and inducible antioxidant defenses in embryos is not well understood. Zebrafish serve as a valuable *in vivo* model to investigate the developmental regulation of the oxidative stress response. The results presented here and in our other recent papers [21,35,46] complement work done previously in this model [32,33,38-40].

### Table 2.

Changes in gene expression in 4-dpf zebrafish eleutheroembryos exposed to tBHQ or TCDD for 6 hr, relative to DMSO-treated eleutheroembryos: Selected genes involved in phase I & phase II biotransformation, GSH synthesis and utilization, oxidative stress response.

| Probe number | Gene | tBHQ mean ± SE | TCDD mean ± SE |
|--------------|------|----------------|---------------|
| A_15_P110246 | catalase | 0.90±0.14 | 0.85±0.03 |
| A_15_P117964 | superoxide dismutase 1, soluble (sod1) | 1.12±0.21 | 0.80±0.07 |
| A_15_P110713 | superoxide dismutase 2, mitochondrial (sod2) | 0.81±0.03 | 0.83±0.04 |
| A_15_P108217 | NADPH dehydrogenase, quinone 1 (nqo1) | 0.66±0.08 | 1.19±0.05 |
| A_15_P107652 | prostaglandin endoperoxide synthase 2 (ptgs) | 5.89±0.48 | 0.61±0.06 |
| A_15_P100578 | cytochrome P450A1 (cytp1a) | 0.96±0.09 | 71.59±6.09 |
| A_15_P102530 | UDP glycosyltransferase (ugt1b5) | 0.66±0.15 | 4.03±0.48 |
| A_15_P101728 | epoxide hydrolase 1, like (epx1l) | 1.03±0.25 | 1.20±0.16 |
| A_15_P100082 | glutamate-cysteine ligase c (gclc) | 2.21±0.12 | 0.97±0.04 |
| A_15_P112437 | glutamate-cysteine ligase m (gclm) | 4.11±0.18 | 1.33±0.09 |
| A_15_P102996 | g-glutamyl transferase (ggta1a) | 7.31±2.65 | 0.84±0.07 |
| A_15_P109364 | glutathione reductase (gr1) | 3.79±0.68 | 1.12±0.04 |
| A_15_P120619 | glutathione peroxidase 4b (gpx) | 0.81±0.06 | 0.89±0.06 |
| A_15_P118489 | glutathione synthase (gss) | 2.19±0.26 | 1.17±0.11 |
| A_15_P112576 | cystathionine beta-synthase (cbsb) | 2.07±0.34 | 0.95±0.07 |
| A_15_P111318 | glutathione S-transferase, alpha-like (gstal) | 1.50±0.27 | 0.68±0.04 |
| A_15_P118878 | glutathione S-transferase, mu (gstm) | 1.14±0.18 | 0.87±0.03 |
| A_15_P107422 | glutathione S-transferase, pi (gstp1) | 2.89±0.65 | 1.50±0.08 |
| A_15_P111132 | microsomal gst 3 | 2.22±0.12 | 0.95±0.08 |
| A_15_P109839 | glutathione S-transferase, omega | 5.25±1.30 | 0.77±0.04 |
| A_15_P103106 | Thioredoxin (txn1) | 5.47±0.92 | 1.04±0.08 |
| A_15_P107290 | similar to vertebrate heme oxygenase decycling 2 (hmo2) | 0.97±0.11 | 1.03±0.13 |
| A_15_P115043 | metallothionein (mt) | 1.07±0.13 | 0.92±0.08 |
| A_15_P105247 | metallothionein 2 (mt2) | 1.92±0.19 | 0.95±0.09 |

Microarray probes with significant (p<0.05, ANOVA with 5% FDR correction) change for tBHQ or TCDD relative to DMSO are in **bold type**. Data represent mean ± standard error of ratio: treated/DMSO.

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Table 3. Changes in gene expression in 4-dpf zebrafish eleutheroembryos exposed to tBHQ or TCDD for 6 hr, relative to DMSO-treated eleutheroembryos: Other stress-responsive genes and transcription factors.

| Probe number | Gene                                              | tBHQ          | TCDD          |
|--------------|---------------------------------------------------|---------------|---------------|
| A_15_P102446 | similar to jun dimerization protein               | 9.21±1.00     | 0.96±0.05     |
| A_15_P117758 | jun B (junb)                                      | 8.71±0.81     | 1.10±0.09     |
| A_15_P101236 | v-jun sarcoma virus 17 oncogene homolog (avian) (jun) | 2.69±0.23     | 0.89±0.07     |
| A_15_P119415 | suppressor of cytokine signaling 3 (socs3)        | 6.45±1.62     | 1.21±0.13     |
| A_15_P113920 | CCAAT/enhancer binding protein (C/EBP), beta (cebpb) | 6.24±1.28     | 0.93±0.04     |
| A_15_P118944 | sprouty (Drosophila) homolog 4 (spry4)            | 5.05±0.85     | 1.19±0.03     |
| A_15_P104781 | v-maf musculoaponeurotic fibrosarcoma oncogene homolog K (avian) (mafK) | 4.08±0.60     | 1.14±0.12     |
| A_15_P102698 | v-maf musculoaponeurotic fibrosarcoma oncogene homolog f (avian) (mafF) | 3.06±0.29     | 1.23±0.13     |
| A_15_P119243 | v-maf musculoaponeurotic fibrosarcoma oncogene homolog family, protein B (avian) (mafB) | 2.04±0.19     | 1.01±0.13     |
| A_15_P101990 | SRY-box containing gene 9a (sox9a)                | 2.44±0.29     | 0.89±0.03     |
| A_15_P109504 | nuclear factor (erythroid-derived 2)-like 2 (nfe2l2; nrf2a) | 1.30±0.04     | 1.19±0.08     |
| A_15_P110831 | nuclear factor, erythroid derived 2, like 1 (nfe2l1; nrf1a) | 0.63±0.21     | 0.72±0.19     |
| A_15_P116909 | nuclear factor, erythroid derived 2, like 1 (nfe2l1; nrf1b) | 2.38±0.51     | 2.19±1.05     |
| A_15_P109440 | nuclear factor, erythroid-derived 2 (nfe2)        | 0.63±0.08     | 0.81±0.15     |
| A_15_P105139 | kelch-like ECH-associated protein 1 (keap1)        | 3.55±0.64     | 1.57±0.23     |
| A_15_P104554 | jun B proto- Oncogene, like (junbl)               | 11.42±1.47     | 1.31±0.03     |
| A_15_P120520 | aryl hydrocarbon receptor 1a (ahr1a)              | 0.64±0.12     | 1.24±0.13     |
| A_15_P103538 | aryl hydrocarbon receptor 2 (ahr2)                | 0.80±0.04     | 2.42±0.24     |
| A_15_P105040 | aryl hydrocarbon receptor nuclear translocator (armt2) | 1.38±0.07     | 1.04±0.04     |
| A_15_P102120 | forkhead box Q1 (foxq1b)                          | 0.43±0.06     | 3.48±0.13     |

Microarray probes with significant (p<0.05, ANOVA with 5% FDR correction) change for tBHQ or TCDD relative to DMSO are in **bold type**. Data represent mean ± standard error of ratio: treated/DMSO.

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43,60,65,66] by expanding the set of known NRF-related proteins potentially involved in regulating the OSR in zebrafish, determining the set of genes that are induced and repressed by a prototypical oxidant (tBHQ), and identifying a phenotype (loss of pigmentation) that is linked to a specific change in gene expression (decrease in mitfa).

Gene expression profiles

An extensive OSR, including both increases and decreases in gene expression, occurred in zebrafish eleutheroembryos exposed to tBHQ for 6 hr at 4 dpf. We were particularly interested in the overlap between the OSR in zebrafish embryos and that described previously in mammalian systems (primarily adult tissues and cells). We also were interested in determining whether there were unexpected changes in gene expression that could represent novel responses to developing vertebrates. Importantly, there was concordance between microarray and qRT-PCR data (Table S5), validating the Agilent zebrafish microarray as a platform for evaluating expression profiles and for gene discovery in developing zebrafish exposed to oxidants. Computational searches for cis-regulatory elements involved in the observed responses to tBHQ and TCDD identified a number of possible regulatory mecha-
Interestingly, in several cases in which there are two or more zebrafish orthologs or co-bryos. More than half of the set of OSR marker genes identified by Johnson and colleagues [47] have zebrafish orthologs such that only one form is inducible, a possible example of subfunction partitioning [43]. In addition to the 10 “biomarker” genes (Table 1), other well-known mammalian OSR genes were induced in zebrafish eleutheroembryos by tBHQ, including several involved in GSH and cysteine homeostasis: glutathione synthetase (cbs), thioredoxin, ferritin heavy polypeptide), only one of the zebrafish ortholog such that only one form is inducible, a possible example of subfunction partitioning [43].

**Overlap in OSR between mammals and developing zebrafish.**

Many, but not all, of the known mammalian oxidant response genes were induced by tBHQ in zebrafish eleutheroembryos. More than half of the set of OSR marker genes identified by Johnson and colleagues [47] have zebrafish orthologs or co-orthologs that were induced by tBHQ in zebrafish at 4 dpf. Interestingly, in several cases in which there are two or more zebrafish co-orthologs of a mammalian gene (hs390, dhnhb1, thioredoxin, ferritin heavy polypeptide), only one of the zebrafish co-orthologs was induced (Table 1), suggesting that the zebrafish paralogs have divided the regulatory features of the mammalian ortholog such that only one form is inducible, a possible example of subfunction partitioning [43].

In addition to the 10 “biomarker” genes (Table 1), other well-known mammalian OSR genes were induced in zebrafish eleutheroembryos by tBHQ, including several involved in GSH and cysteine homeostasis: *gcl*, *gr1*, glutathione reductase (*gr2*), glutathione synthetase (*gss*), thioredoxin (*thd1*), cystathionine beta-synthase (*cbs*), and two GSTs (*gsta* and *gste*). We also saw strong induction of gamma-glutamyltransferase (*ggt1a*), which is involved in an extracellular GSH salvage pathway [67] and is important in protecting against...
oxidative DNA damage [68]. Other genes induced by tBHQ in our study that have also been reported to respond to oxidative stress in at least some mammalian systems include *atf3* and *dusp1* [47] (Table 3).

Overlap in OSR among studies in zebrafish embryonic stages and adults. Some of the OSR genes induced in our study (e.g. *txn1*, *gstp*, *gsto1* *hsp70*, *dnaj*, *atf3*) were also shown to be induced by tBHQ in a study comparing the response of zebrafish embryos to several different toxicants, using a different and less complete microarray platform, published while this manuscript was in preparation [60]. Tanguay and colleagues [69] reported that fullerene (C60) caused an OSR in zebrafish embryos; genes induced in common by C60 [69] and tBHQ (this study) include *hsp70*, *gstp1*, *gclc*, and ferritin. There are no published studies showing effects of tBHQ on adult zebrafish. However, the response of adult zebrafish liver to arsenic (Na2HAsO4) exposure included genes associated with an OSR, including several that were also induced by tBHQ in our study (*hsp70*, *hsp90a*, ferritin, *gstp1*, *gsto1*, *txnrd1*, *txn*, *gadd45b*). Arsenic also induced hepatic expression of *sod2*, *gpx4b*, and *mt2*, which were not induced by tBHQ in embryos under the conditions examined in our experiment, demonstrating compound-, concentration-, tissue-, or stage-specific differences in the response to oxidants.

There are several other interesting differences in our data as compared to results obtained in other systems. Recent studies suggest that tBHQ is a weak AHR agonist in mammals [70,71], but at the concentration used in our experiments (10 μM) tBHQ did not activate the AHR, as indicated by the absence of *cyp1a* expression.

Figure 4. Changes in gene expression in developing zebrafish following exposure to TCDD (2 nM) or tBHQ (10 μM) in 0.1% DMSO. Embryos, eleutheroembryos, or larvae at 1, 2, 3, 4, 5, or 6-dpf were exposed to each chemical for 6 hours, after which they were frozen for RNA isolation and analysis of gene expression. Expression of *hsp70*, *gadd45b*, *atf3*, *mitfa*, *opn1lw1*, and *foxq1b* genes were measured by qRT-PCR. Values represent mean±SE of 4 biological replicates, each replicate a pool of 30 embryos. *statistical significance at p<0.05 (Dunnett’s test).

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Figure 5. Phenotypic changes following tBHQ exposure. Zebrafish embryos were exposed to tBHQ (5 or 10 μM) or 0.1% DMSO from 32–47 hpf. A. At 52 hpf, randomly selected embryos were mounted in 3% methylcellulose and imaged as described in Materials and Methods. B. Embryos were subsequently maintained in 0.3× Danieu’s water (without tBHQ) at until 120 hpf, when they were imaged again in order to assess recovery of pigmentation.

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Figure 6. EGFP-HSP70 zebrafish exposed to tBHQ. A. Embryos from hsp70-EGFP transgenic zebrafish were exposed to DMSO (0.1%) or tBHQ (10 μM) for 4 hrs at 1-dpf. After exposure, embryos were washed and held for 4 additional hours prior to fluorescence microscopy for detecting EGFP expression. B. Another experiment was performed by exposing 4-dpf eleutheroembryos (n=15) to DMSO (0.1%) or tBHQ (10 μM) for 4 hrs at 28°C. After exposure, eleutheroembryos were washed with 0.3× Danieu’s and inspected 4 hrs post-exposure for EGFP expression by fluorescence microscopy.

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induction in developing zebrafish at 1–6 dpf and by the negligible overlap between tBHQ- and TCDD-modulated gene sets.

Another notable aspect of our results is the genes not significantly affected by tBHQ exposure. For example, in contrast to what is observed in mammals [18,47], there was no induction of nqo1 or sod1 by tBHQ as assessed either by microarray (in the 4-dpf eleutheroembryos) or by qRT-PCR (all time points). The lack of nqo1 induction differs from the results of Kobayashi [33], who reported induction of this gene in 4-dpf zebrafish larvae exposed to 30 μM tBHQ for 6 hr. This could reflect a difference in the concentrations used in the two studies (10 μM vs. 30 μM). The induction of sod genes (sod1 and sod2) in zebrafish embryos may be compound-specific, as suggested by the results of Timme-Laragy et al. [39], who observed induction of sod1 and sod2 after exposure to a mixture of flavonoids but not after exposure to tert-butyldihydroperoxide (at a concentration that induced other OSR genes such as gce1 and gstp1). Despite the lack of induction of nqo1 or sod1 observed in our experiments, there may be other exposure conditions under which tBHQ might induce these genes.

Novel and notable responses. The microarray results also revealed novel changes in gene expression. We mention just a few of these to illustrate the apparent richness of the oxidative stress response in zebrafish eleutheroembryos. Hypoxia-induced gene (hig1) was induced 7-fold by tBHQ. This gene, first identified in fish exposed to hypoxia [72], is closely related to human hspc010 (hematopoietic stem-cell progenitor cell gene 10) [73]. tBHQ also strongly induced prostaglandin endoperoxide synthase 2 (pge2: cog2), dual-specificity phosphatases (dusp5, dusp4, and dusp1), insulin-like growth factor binding protein (igfbp1), and hepcidin anti-microbial peptide (hamp). These and other changes suggest the response to tBHQ in developing zebrafish involves, in addition to a classical OSR, transcriptional changes resembling an inflammatory response (pge2), response to hypoxia (hig1, igfbp1), and responses to maintain iron homeostasis (hamp, ferritin).

In tBHQ-treated eleutheroembryos at 4 dpf there was strong induction of genes associated with a DNA-damage response, such as atf3 [74], gadd15 (several forms), and dnanf1 (both, suggesting that oxidative DNA damage may be an early effect of this compound. We showed recently that the induction of atf3 by tBHQ in zebrafish embryos was not controlled by Nrf2a or Nrf2b [21], consistent with the idea that at least some of these responses may be secondary to damage rather than a direct effect of tBHQ mediated through one of the Nrf2 proteins.

The results also revealed additional induced genes of interest. For example, several zebrafish members of the solute carrier family (e.g. slc25a25, slc13a2, slc16a9a, slc16a3, slc1a4, slc16a6b) were induced by tBHQ. Human slc25a25 encodes a mitochondrial carrier that transports adenine nucleotides (ATP) across the inner mitochondrial membrane in exchange for phosphate [75]. Three monocarboxylate transporters (MCTs) (slc16a6b, slc16a9a, and slc16a3) were induced ~3–5-fold; MCTs transport pyruvate, which can scavenge ROS [76]. SLC genes have not been generally recognized as part of the oxidative stress response. However, SLC3A2 was recently shown to be induced in human umbilical vein endothelial cells exposed to lipid oxidation products derived from oxidized low-density lipoproteins [77]. SLC3A2 and SLC1A1 (amino acid transporters) were induced in HepG2/C3A cells in response to cysteine deprivation [78], and SLC1A1 (a cysteine transporter) was induced by tBHQ and sulfurophane in rat glioma cells [79]. Although slce3a2 and slce1a were not among the slc genes induced in our experiment, overall the increase in slc gene expression suggests that induction of certain transport proteins might be an important part of the oxidative stress response, or a wider "integrated stress response" [80] in developing vertebrates. Induction of sde genes also occurs in response to other toxicants in zebrafish embryonic stages, but the patterns of sde gene expression may be toxicant-specific [60].

Implications for response to other oxidants. tBHQ is widely used as a prototypical mono-functional inducer of the OSR [23], but whether the results obtained here are representative of changes expected from exposure of developing zebrafish to other antioxidant chemicals is not clear. The chemical specificity of the OSR has not been thoroughly investigated, especially during vertebrate development. Changes in gene expression (gene expression profiles) may vary according to the type or localization of oxidative stress, for example as caused by different types of pro-oxidant chemicals or other mechanisms of oxidative stress. Although some studies report similar expression profiles in response to different types of oxidative stress [81,82], there is evidence for chemical-specific roles of Keap1 paralogs and mechanistically distinct classes of Nrf2 activators [39] and there are reports of distinct gene expression patterns from genetic versus chemical activation of Nrf2 [83,84]. Our recent studies show that Nrf2a and Nrf2b regulate distinct but partially overlapping sets of genes constitutively [21] and in response to tBHQ (manuscript in preparation). Thus, it will be important to determine how embryos respond to chemicals that generate oxidative stress via different mechanisms. Such studies are underway [85].

Response to TCDD. One goal of this work was to compare the transcriptional response of developing zebrafish to TCDD exposure with that caused by a model oxidant such as tBHQ. Oxidative stress has been implicated in effects of TCDD [86–88] including in embryonic stages [52], but after 6 hr of exposure to TCDD at 4 dpf we found no evidence for an oxidative stress response in the genes measured by microarray. Similarly, targeted analysis of gene expression by qRT-PCR at all six exposure times (days 1 through 6) showed no changes in classical OSR genes such as gce1, gce, or sod1 (Fig. 2). Other investigators also found little evidence for an OSR in developing zebrafish exposed to TCDD [89]. Our experiments in whole animals (embryos, eleutheroembryos, and larvae) could have missed highly localized oxidative stress and resultant changes in gene expression. We did, however, observe induction of gce1 in whole embryos 48 hr after exposure to TCDD starting at 24 hpf (unpublished studies), suggesting that generation of oxidative stress or disruption of sulfhydryl balance may be delayed or may occur in embryos after prolonged exposure to this compound. Though widespread oxidative stress does not appear to be part of the acute response to TCDD exposure, it may occur in specific cell types or be part of the response to longer-term exposure to TCDD.

One of the more interesting changes observed in TCDD-exposed embryos was the induction of a foxq1 gene now called foxq1b. After our preliminary reports of these data [90,91] and while this manuscript was in preparation, Planchart & Mattingly [92] reported the induction of a different foxq1 gene (now called foxq1a) by TCDD in zebrafish embryos. The TCDD-induced foxq1 gene reported by Planchart & Mattingly [92] was called foxq1b in that paper but now has been renamed foxq1. It is located on zebrafish chromosome 2, encodes predicted protein XM_003197808.1, and corresponds to Agilent probe A_15_P102120. We will use the currently approved nomenclature foxq1a (the gene reported induced by Planchart & Mattingly) and foxq1b (the gene reported to be induced in the current study).
In the Planchet & Mattingly study [92], foxq1b (new nomenclature) was not induced by TCDD (1 nM). However, those authors examined embryos at 24 and 48-hpf. In our studies, foxq1b was also not inducible by TCDD (2 nM) at those early times, but became highly inducible at later times (3, 4, and 5 dpf) [Fig. 4], demonstrating stage-specific responsiveness not seen with foxq1a [92]. The murine Foxl gene also appears to be responsive to TCDD [93], evidence for an evolutionarily conserved role of AHR in regulating foxl genes. Although the zebrafish foxl gene is expressed in jaw primordia, the site of foxl expression is not yet known and the functions of these two paralogs in zebrafish development have not yet been investigated. In mammals, FOXL1 is expressed both in embryos and a variety of adult tissues [94,95], is required for normal embryonic development [94,96], and has a recently discovered role in controlling epithelial-mesenchymal transition in human cancer metastasis [97-99]. It will be important to better characterize the relationship between foxl and ahr genes and their roles in cellular and developmental processes.

Ontogeny of antioxidant response

Although it is known that the sensitivity of developing vertebrate animals to chemicals varies by developmental stage [60,100-102], the underlying mechanisms are not well understood. Previous research in a variety of vertebrate models has suggested that in early development the antioxidant defense systems are immature and not fully responsive to oxidative stress [33,103,104]. However, we are not aware of any systematic investigations of the developmental stages at which vertebrates develop the capacity to respond to oxidative stress by the induction of anti-oxidant defenses. Here, we found that embryos as early as 24 hpf were capable of responding to tBHQ with induction of gsp1, gclc, and nrf2a. The result with gsp1 differs from that of Kobayashi et al. [33], who found by in situ hybridization that expression of gsp1 was inducible at 96 and 120 hpf, but not at 24 hpf. Our results suggest that qRT-PCR is more sensitive for detecting induced gsp1 at this early time.

Our results also showed that the response to tBHQ varied by developmental time, in a gene-specific manner. For example, gsp1 and gclc, while inducible at 1- and 2-dpf, were less inducible (and more highly variable) at later stages. These results suggest that the set of genes responsive to tBHQ or other oxidants will vary as embryos and later stages develop. This could be related to developmentally programmed changes in GSH reductase status [35]. In future studies, it will be important to examine the relationships among GSH status, inducibility of antioxidant defenses, and stage-specific differences in sensitivity to embroytotoxicity of oxidant chemicals. Such studies should include an assessment of dose-response relationships and how they may change during development.

Repression of mitf and pigmentation defects caused by tBHQ

One of the genes that we found by microarray (Table 4) and qRT-PCR (Fig. 4) to be most strongly repressed in tBHQ-treated embryos and eleutheroembryos was mitfa, one of two zebrafish orthologs of the mammalian gene MITF [61]. Expression of MITF has also been reported to be reduced in mammalian melanocytes exposed to 4-tert-butyphenol [105] or melanoma cells exposed to hydrogen peroxide [106]. MITF and mitfa control melanocyte differentiation and regulate the expression of enzymes involved in melanin synthesis [107]. mitfa is defective in zebrafish nacre mutants, which exhibit hypopigmentation [62], and knockdown of Mifa protein in zebrafish embryos with morpholino-modified antisense oligonucleotides causes nearly complete but transient loss of body pigmentation [108,109]. We found that zebrafish embryos exposed to tBHQ exhibited reduced pigmentation, confirming and extending a previous report [60]. Partial recovery of pigmentation occurred several days after cessation of tBHQ exposure. These results provide a link between oxidative stress, mitfa, and loss of pigmentation, and suggest that tBHQ-treated zebrafish embryos could serve as a model for vitiligo, a human skin disease characterized by depigmentation and reduced expression of MITF in melanocytes [110]. Vitiligo has been suggested to have an etiology involving oxidative stress [111] and NRF2 polymorphisms were identified as risk factors in the development of this disease [112].

hsp70-GFP transgenic zebrafish as tool for screening

Our microarray and qRT-PCR data showed that hsp70 was induced at all stages and up to 50-fold in zebrafish early life stages exposed to tBHQ (Table 3, Fig. 4). Consistent with this, we found two putative AREs in the promoter of the zebrafish hsp70 gene, although they were identified only when using a high false discovery rate. The strong response of hsp70 to tBHQ treatment prompted us to evaluate the effect of tBHQ on embryos of an hsp70-egfp transgenic zebrafish line [33], which has been shown previously to respond to heat or cadmium with induced expression of the egfp transgene [33,113]. We found that tBHQ exposure early in development caused induction of EGFP in a restricted pattern, whereas exposure later in development caused widespread EGFP expression (Fig. 6).

The widespread induction of hsp70-egfp at 4-dpf is consistent with the high level of induction of hsp70 measured by microarray and qRT-PCR and with studies in human cells showing that HSP70 is regulated by NRF2 [63]. The strong response of hsp70, hsp40, jun, and fos could reflect a direct effect of signaling via ROS, potentiated by GSH depletion [114]. These results demonstrate the potential utility of EGFP transgenic fish for assessing cell- and tissue-specific effects of oxidant chemicals, complementing whole-embryo assessments of gene expression by qRT-PCR and microarray and providing a method for rapidly screening chemicals for the ability to cause oxidative stress during development. However, the hsp70-egfp fish respond to a variety of stressors [53,113,115], and thus lack the specificity that would be required for a targeted screening assay. Thus, it will be important to develop germ-line transgenic fish lines expressing reporter genes under control of more specific indicators of oxidative stress [43,65,66]. The gene expression data reported here will help to identify the appropriate target genes as a source of regulatory elements for use in such an approach.

Conclusions

The key findings of these experiments are: 1) embryos are responsive to tBHQ as early as 24-hpf, with strong induction of classical OSR genes like gsp1 and gclc; 2) the response to tBHQ varies with developmental time, in a gene-specific manner; 3) at 4-dpf, tBHQ induces a suite of OSR genes including several involved in GSH metabolism, response to DNA damage, amino acid transport, response to hypoxia, iron homeostasis, and inflammation; 4) microarrays were capable of detecting altered expression of a variety of known and novel oxidant-responsive genes in whole eleutheroembryos; and 5) patterns of tBHQ-induced gene expression in developing zebrafish exhibit strong similarities but also some differences as compared with genes induced by tBHQ in mammalian systems (adults and cultured cells). These data demonstrate the responsiveness of developing zebrafish Response to tBHQ and TCDD during Development
zebrafish to a model oxidant (tBHQ) and illustrate the power of this approach for investigating the mechanisms by which early life stages of vertebrate animals respond to oxidative stress. The results will help guide studies using zebrafish embryonic and larval stages to better understand the chemical and stage specificity of the OSR and its role in determining the sensitivity of vertebrate animals to oxidant chemicals during development.

Materials and Methods

Chemicals

2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD) was obtained from Ultra Scientific (Hope, RI), Dimethyl sulfoxide (DMSO) was from Sigma-Aldrich (St. Louis, MO), Tert-butylhydroquinone (tBHQ) was obtained from Acros Organics (Geel, Belgium).

Embryo culture

For experiments 1 and 2, we used adult zebrafish of the TL strain, a generous gift of Dr. Mark Fishman (Massachusetts General Hospital, Cambridge, MA), to generate embryos. Fish were maintained as described previously [116]. For experiment 3, we used hsp70-EGFP fish [Tg(hsp70:EGFP)_unspecified] [53], a generous gift from Dr. John Y. Kuwada (University of Michigan).

The experiments were carried out according to the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The protocols were approved by the Institutional Animal Care and Use Committee of the Woods Hole Oceanographic Institution (IACUC Assurance: A3630-01).

Exposure of embryos, eleutheroembryos, or larvae to chemicals

Experiment 1. Separate groups of 30 embryos generated from TL adults were placed in 20 ml system water in 10-cm glass petri dishes. At 1, 2, 3, 4, 5, and 6-dpf, embryos, eleutheroembryos, or larvae were exposed for 6 hr to DMSO (0.1%), TCDD (2 nM), or tBHQ (10 μM) (4 groups per compound at time point) (Fig. 1). The concentration of TCDD (2 nM) is known to strongly induce expression of gene development in developing zebrafish [51,60]. The concentration of tBHQ (10 μM) is one at which strong induction ofgstp1 was reported in 4-dpf eleutheroembryos after 6 hr exposure [33]; higher concentrations (30 μM) caused acute toxicity in our hands. Embryos were frozen immediately after the 6-hr exposure. The short exposure time was chosen to select for primary (direct) responses and minimize changes in gene expression that are secondary to toxic effects. RNA was isolated for real-time, quantitative reverse-transcription-polymerase chain reaction (qRT-PCR) and microarray analysis, as described below. (Throughout the paper, we refer to these embryos by the age when they were inspected and imaged for pigmentation. After exposure, eleutheroembryos were washed with 0.3× Danieau’s and its role in determining the sensitivity of vertebrate animals to oxidant chemicals during development.

Experiment 2. Groups of 150 TL embryos were exposed to tBHQ (5 or 10 μM) or 0.1% DMSO in glass petri dishes containing 20 ml of 0.3× Danieau’s from 32–47 hpf, then washed in 0.3× Danieau’s at 52 hpf, randomly selected embryos were mounted in a left-lateral orientation in 3% methylcellulose and imaged using a Zeiss axioimager using a Zeiss Axiocam MR color CCD camera. Embryos were subsequently maintained in 0.3× Danieau’s at 28.5 °C until 120 hpf, when they were imaged again in order to assess recovery of pigmentation.

Experiment 3. Separate groups of 1-dpf embryos (n = 30) from hsp70-EGFP transgenic fish were exposed to DMSO (0.1%) or tBHQ (10 μM) for 4 hrs at 28°C. After exposure, embryos were washed and held for 4 additional hours in 0.3× Danieau’s before being inspected for EGFP expression by fluorescence microscopy using an Axiovert 200 inverted microscope with Zeiss Filter Set 38 HE (409038; excitation BP 470/40, FT 495, emission BP 525/50). Another experiment was performed by exposing 4-dpf eleutheroembryos (n = 15) to DMSO (0.1%) or tBHQ (10 μM) for 4 hrs at 28°C. After exposure, eleutheroembryos were washed with 0.3× Danieau’s and its role in determining the sensitivity of vertebrate animals to oxidant chemicals during development.

RNA isolation and qRT-PCR

Total RNA was isolated using RNA STAT-60 (Tel-Test B, Inc., Friendswood, TX) and DNase-treated using the Turbo DNA-free kit (Ambion, Austin, TX). Poly(A)+ RNA was purified using the MicroPoly(A)Purist Kit (Ambion). cDNA was synthesized from 2 μg of total RNA using Omniscript reverse transcribe transcriptase (Qiagen, Valencia, CA). Real-time qRT-PCR was performed using the iQ SYBR Green Supermix (Bio-Rad, Hercules, CA) in an iCycler IQ Real-Time PCR Detection System (Bio-Rad) as described previously [117]. Primers were synthesized by Midland Certified Reagent Company, Midland, TX. Primer sequences can be found in Table S1.

Microarray analyses

We examined a subset of RNA samples from Experiment 1 by microarray using the Agilent 22k long-oligo zebrafish array. We analyzed all of the 4-dpf samples: four biological replicates each for DMSO-, tBHQ- and TCDD-treated eleutheroembryos, each hybridized against a universal reference mRNA created from equal amounts of RNA from 2 replicates each from all toxicants (TCDD, tBHQ, DMSO) and time points (1, 2, 3, 4, 5, 6-dpf). The use of a universal reference RNA balances efficiency with statistical power [118] and has several advantages [119-121]. It facilitates normalization because all of the genes expressed in experimental samples are represented in the reference samples [122]. Dye bias is minimized because all experimental samples are labeled with the same dye; thus, dye swaps are not needed [121]. To verify this, we performed quality control hybridizations including a dye swap and a self-self hybridization. Analysis of a self-self hybridization of the Universal Reference (composed of equal amounts of RNA from all timepoints, toxicants, and replicates) revealed 18293 of 21495 features (85%) with signal above background (calculated as 2.6 times the background standard deviation). This indicates that the majority of the probes on the Agilent microarray represent transcripts expressed in 1–6 dpf embryos.

RNA samples from 4-dpf eleutheroembryos treated with DMSO, tBHQ, or TCDD were checked for quality using a NanoDrop ND-1000 spectrophotometer and an Agilent 2100 Bioanalyzer. cDNA synthesis from 200 ng of total RNA was performed using the Agilent Low RNA Input Linear Fluorescent Amplification Plus kit, following the manufacturer’s instructions. cRNA was synthesized from the cDNA template, with incorporation of either cyanine-5-CTP or cyanine-3-CTP (Perkin Elmer). Labeled cRNA was purified using the Qiagen RNeasy Mini kit; quantity and quality was assessed by NanoDrop spectrophotometer and Agilent Bioanalyzer. cRNA samples were hybridized to Agilent 22k zebrafish microarrays using the Agilent Gene Expression Hybridization Kit. An aliquot (750 ng) of each cy3- and cy5-labeled sample cRNA was hybridized against 750 ng of cy3- and cy5-labeled cRNA derived from the Universal mRNA Reference. Labeled cRNAs were combined with the Agilent 25× fragmentation buffer and incubated at 60°C for 30 minutes. This was followed by mixing with 2× hybridization buffer, after which 100 μl of the product was spread evenly across the surface of an
Agilent 22K zebrafish microarray. The loaded microarray was incubated at 60°C for 17 hours with rotation in an Agilent DNA Microarray Hybridization Oven. Post-hybridization, microarray slides were washed, air-dried, and stored in darkness with desiccation prior to laser-excited fluorescence scanning in an Agilent DNA Microarray Scanner.

Analysis of raw microarray data was performed using Agilent’s feature extraction protocol, which includes spot finding, spot analysis, background subtraction (using local background plus global background based on spots along the central tendency line for red versus green intensity), dye normalization (linear and lowess algorithms, using spots along the central tendency line as for background subtraction), and final calculation of Cy5/Cy3 ratios and log₂ transformed fold change for each spot. Features with signal not significantly above background, non-uniform features, and features exhibiting saturation were flagged. The microarray data have been deposited in MIAME-compliant format in the Gene Expression Omnibus (GEO) database at the U.S. National Center for Biotechnology Information (accession number GSE10157; http://www.ncbi.nlm.nih.gov/geo/).

Probes indicating significantly different relative transcript abundance among the DMSO, TCDD, and tBHQ treatments were determined by Analysis of Variance (ANOVA) using MEV in the TM4 suite of microarray software [123,124]. Data were first log transformed and values for each probe median centered. ANOVA was performed with a distribution based on 1000 permutations of the data, a significance value of p<0.05, and control of False Discovery Rate (FDR) at 5% [125,126]. An objective of the statistical analysis was to minimize type II error while maintaining a reasonable false discovery rate.

Hierarchical cluster analysis of the significant probes was performed using CLUSTER software [127] and log₂ transformed fold change values were median centered for both probes and microarrays. Cluster analysis used Pearson’s correlation (i.e. centered) and average linkage clustering. Enrichment of Gene Ontology (GO) terms for clusters of significant probes thus identified was examined using the FatiGO+ software [56]. FatiGO+ uses GO terms assigned within the Ensembl annotation of the zebrafish genome [128] and the background set of probes used in each analysis was the entire probe set of the Agilent microarray less the probes found significant by the ANOVA analysis described above.

Annotation of probes on the array

To aid interpretation of all results, we updated the microarray annotation provided by Agilent by incorporating annotations, functional domain predictions, and Gene Ontology assignments available at Ensembl (version 48, based on assembly Zv7), ZFIN, UniProt, ReSeq, and the Harvard Gene Index Annotation. Additional putative annotations were obtained from Ensembl’s Integr8 project [129], which attempts annotation of proteins based upon putative orthology among organisms. Comparison to the latest Zv9 assembly assignment of Agilent probes to genes at Ensembl did not improve gene annotations.

A number of probes found important in our statistical analyses were manually annotated by searching the zebrafish ReSeq RNA database or available zebrafish ESTs for the putative transcript and examining the similarity of the encoded protein to those of other model organisms by using BLAST against the GenBank non-redundant protein database.

**Prediction of cis-regulatory elements**

The 10 kb upstream regions of all genes predicted by Ensembl (version 59, genome assembly Zv9) were searched for putative cis-regulatory motifs using the FIMO software [130]. Searches were for motif matches in either orientation/strand in the target upstream region and were based on position-specific scoring matrix (PSSM) searches with p<0.0001 and associated calculation of q-values [131], with correction for background nucleotide frequencies of complete nuclear genome sequences. Background frequencies were estimated by examining the complete nuclear chromosome data of the Zv9 assembly. Searches used PSSM models for ARE [132], DRE [133], ERE [134], NRF2 [135], PXR [136] and a subset of the JASPAR database [137]: Arnt, Arnt::Ahr, Nf-kappaB, Sp1, Nfe2L1::MafG, Ap1, Rel, NfkB1, RelA, MafB, Nfe2L2, Hif1A::Arnt. We focused our attention on significant hits (p<0.0001) with false-discovery rate of 10% or less.

**Supporting Information**

Table S1 Primers for Real-time RT-PCR. (DOCX)

Table S2 List of all significant probes and the ratios (tBHQ/DMSO, TCDD/DMSO). (XLS)

Table S3 Enrichment of Gene Ontology (GO) terms for Ensembl annotated probes with significantly different abundance between TCDD, tBHQ, and DMSO treated eleutheroembryos. (DOC)

Table S4 List of 2-fold significant probes. (XLS)

Table S5 Comparison of altered gene expression as measured by microarray and qRT-PCR. (DOC)

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**Author Contributions**

Conceived and designed the experiments: MEH SIK DGF AGM [JS BRW EL. Evaluated the experiments: SIK DGF BRW MJJ ART-L AGM MJC. Performed the experiments: SIK DGF BRW MJJ ART-L AGM MJC. Analyzed the data: AGM MJC RIH SIK DGF MJJ ART-L. Contributed reagents/materials/analysis tools: EL. Wrote the paper: MEH AGM. Reviewed and edited the manuscript: SIK DGF AGM MJJ ART-L, [JS BRW MJC EL.

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