Monoamine oxidase A (MAO-A) and MAO-B metabolize biogenic amines and have been implicated in neuronal apoptosis. Although apoptosis is an important process in embryo development, the role of MAO isoforms has not been investigated in detail. We found that expression of MAO-A and MAO-B can be detected early on during embryo development. Expression levels remained constant until around midgestation but then dropped to almost undetectable levels toward birth. Similar expression kinetics were observed in the brain. Isoform-specific expression silencing of MAO-A mediated by siRNA during in vitro embryogenesis induced developmental defects, as indicated by a reduction of the crown rump length and impaired cerebral development. These alterations were paralleled by elevated serotonin levels. Similar abnormalities were observed when embryos were cultured in the presence of the MAO-A inhibitor clorgyline or when the transcriptional inhibitor of MAO-A expression R1 was overexpressed. In contrast, no such alterations were detected when expression of MAO-B was knocked down. To explore the underlying mechanisms for the developmental abnormalities in MAO-A knockdown embryos, we quantified the degree of developmental apoptosis in the developing brain. MAO-A knockdown reduced the number of apoptotic cells in the neuroepithelium, which coincided with impaired activation of caspases 3 and 9. Moreover, we observed apoptotic cells in the neuroepithelium, which coincided with reduced cyclin D1 levels as an indicator of impaired cell proliferation. This data highlights MAO-A as a vital regulator of embryonic brain development.

Monoamine oxidases (MAO-A, MAO-B) metabolize biogenic amines and have been implicated in neuronal apoptosis. This reaction requires molecular dioxygen and produces stoichiometric amounts of hydrogen peroxide and ammonia. In higher animals, there are two MAO isoforms, referred to as MAO-A and MAO-B, which share a high degree (70%) of amino acid conservation. The two isoforms can be distinguished with respect to their metabolism of xenobiotics. MAO-A is predominantly found in catecholaminergic neurons, and highest concentrations are detected in the locus coeruleus. In contrast, MAO-B is most abundant in serotonergic and histaminergic neurons as well as in glial cells. In the periphery, distribution of MAO isoforms is variable, and there are species-specific differences.

The two MAO isoforms are encoded for by separate genes on the X chromosome. The promoter regions of the genes have been investigated, and the transcriptional repressor R1 was found to specifically bind and repress the MAO-A promoter. A functional promoter polymorphism of the MAO-A gene has been related to behavioral disorders. In humans, functional inactivation of MAO genes is associated with a severe form of Norrie disease. In addition, pharmacologic inactivation of MAO activity is a therapeutic approach for the treatment of neurological disorders that have been shown to involve MAO activity, such as depression or neurodegenerative disorders, including Parkinson’s disease and Alzheimer’s disease.

To study the in vivo role of MAO isoforms in cerebral function, MAO-deficient mice have been employed. Targeted knockout of the MAO-B gene leads to viable offspring that show an altered stress response. Unfortunately, to date targeted inactivation of the MAO-A gene has not been carried out. However, naturally occurring MAO-A-deficient mouse strains have been identified. Incidental insertion of an interferon β minigene in exon 2 of the MAO-A gene abrogates the expression of a functional enzyme. These mice are viable and do not show obvious signs of functional defects in a resting state. However, they have significantly increased plasma levels of monoaminergic transmitters (serotonin, dopamine, norepinephrine), are more susceptible to different kinds of stress, and exhibit a tendency for aggressive behavior. Similar observations were made in mice bearing a spontaneous point mutation in exon 8, which introduced a premature stop codon analogous to a mutation found in humans suffering from Brunner
syndrome (14). An identical mutation occurred in a colony of MAO-B-deficient mice, leading to MAO-A/B double knockout animals that are also viable but exhibit elevated monoamine levels and anxiety-like behavior (15).

Monoamines are essential for proper cerebral function. Monoaminergic transmitter systems appear early on in embryogenesis (16), and serotonin, the biologically most relevant MAO-A substrate, has been implicated in maturation of neuronal progenitors (17). MAO-A-deficient mice lack developmental clustering of layer IV granular neurons and suffer from aberrant maturation of the brainstem respiratory network (18, 19). Although the underlying mechanisms have not been studied in detail, these effects could be explained by the inability of the animals to metabolize serotonin (20). This data is consistent with the recent observation that transgenic overexpression of MAO-A in the forebrain of MAO-A knockout mice rescues their aggressive behavior (21).

Because the outcome of functional studies employing systemic stem cell knockouts is frequently impacted by genetic drifts and epigenetic phenomena (22, 23), we employed the siRNA technology to explore the role of MAO isoforms in embryogenesis. This experimental setup allows us to study the impact of MAO expression at greater spatial and temporal specificity. For this purpose, we explanted murine embryos at early developmental stages, injected siRNA probes or specific inhibitors into the amniotic cavity, and cultured the treated embryos for up to 72 h in vitro (24). Using these genetic and pharmacological intervention strategies, we found that knockdown of MAO-A expression disturbed embryonic brain development and dysregulated developmental apoptosis.

**MATERIALS AND METHODS**

**Chemicals**—The chemicals used were from the following sources: Superscript III reverse transcriptase and RNaseOUT from Invitrogen, BD Advantage 2 polymerase mix from BD Biosciences, dNTPs from Carl Roth GmbH (Karlsruhe, Germany), and PCR primers from BioTeZ Berlin-Buch GmbH (Berlin, Germany).

**RNA Extraction and Reverse Transcription**—Total RNA was extracted using the RNeasy mini kit (Qiagen, Germany) and was reversely transcribed according to standard protocols with oligo d(T)15 primers and SuperScript III reverse transcriptase (Invitrogen) according to the vendor’s instructions.

**Quantitative RT-qPCR**—RT-qPCR was carried out with a RotorGene 3000 system (Corbett Research, Australia) using ImmoMix/SYBR Green (Bioline, Germany). Isoform-specific amplification primers were designed, and their sequences are given in supplemental Table S1. Absence of cross-amplification between the two isoforms was ensured, and the standard protocol has been described before (24). RNA preparations were analyzed at least in triplicate, and mean values are given. The experimental raw data were evaluated with the RotorGene Monitor software (version 4.6). To generate standard curves for quantification of expression levels, specific amplicons were used as external standards for each target gene (5 × 10^3 to 3 × 10^9 copy numbers). GAPDH mRNA was used as the internal standard to normalize expression of the target transcripts.

**MAO Knockdown and Embryogenesis**

Preparation of Mouse Embryos and in Vitro Culture—All animal experiments were performed in strict adherence to the guidelines for experimentation with laboratory animals set in institutions. Ibmred Institute for Cancer Research pregnant mice were obtained from the animal house, and embryos at different developmental stages (gestational day 6.5 (E6.5) to E17.5) were prepared under a stereomicroscope (Olympus). For RT-qPCR, preparations were kept in PBS (0.1% diethyl pyrocarbonate), and extraembryonic tissue was removed. For in vitro culture, the embryos were dissected in modified phosphate-buffered saline medium (5% FBS) as described before (24).

Transfection Experiments—For our knockdown strategy, we designed isoform-specific siRNA using the Stealth™ RNAi program (BLOCK-iTTM RNAi Designer, Invitrogen). Among the designed probes we selected those constructs supplemental Table S1 exhibiting highest estimated knockdown probability and a GC content below 40%. No sequence cross-homology was allowed. Control siRNA duplexes with no homology to any vertebrate transcript were used as random control probes (Stealth™ RNAi negative controls, Invitrogen). For R1 overexpression, the R1 coding sequence was cloned into the mamma- lian pcDNA3.1 expression vector supplemental Table S1). Empty vector was used as a control. For knockdown experiments, murine embryos were explanted at E7.5 and transfected with siRNA constructs. 10 nl of annealed double-stranded siRNA (25–100 nM) were mixed with 0.01% Lipofectamine™ 2000 (Invitrogen) and then microinjected with an Application Solution Transgenic platform micromanipulator (Leica, Germany) into the amniotic cavity. After microinjection the embryos were placed in a whole embryo culture roller incubator (BTC Engineering, UK). The embryos at early gastrulation stage (E7.5) were allowed to develop for up to 72 h until early organogenesis stages E9.5–E10.5 in 100% heat-inactivated rat serum with a continuous flow of gas mixtures (24).

Evaluation of Developmental Milestones—The impact of the treatments on embryonic growth (embryo size measured by the crown rump and head length) and on development of the brain, heart, and limbs was quantified by a microscopic scoring procedure. To judge cerebral embryogenesis, we assessed the degree of brain maturation by a number of morphological parameters (25). A score of 0 represented strong developmental retardations, whereas a score of 5 indicated normal development. The following morphological criteria were applied for the different parts of embryonic brain: Forebrain: prosencephalon invisible (score 0), V-shape (score 1), U-shape (score 2), partially fused (score 3), completely fused (score 4), telencephalic evaginations (score 5). Midbrain: mesencephalic invisible (score 0), V shape (score 1), U shape (score 2), partially fused (score 3), completely fused (score 4), discernible division between mesencephalon and diencephalon (score 5). Hindbrain: rhombencephalon invisible (score 0), V shape (score 1), U shape (score 2), folds fused with anterior neuropore (score 3), anterior neuropore closure (score 4), dorsal flexion develops with 4th ventricle roof (score 5). Significances were calculated with Students’ t test.

MAO Activity Assays—Enzymatic activity of MAO isoforms was assayed in vitro with 14C-labeled tyramine hydrochloride as
a substrate, as described previously (26). The relative share contributed by the two MAO isoforms was quantified using isoform-specific irreversible inhibitors. Clorgyline was used for specific inhibition of MAO-A and deprenyl for inhibition of MAO-B. In Situ Hybridization—In situ hybridization was performed according to Wilkinson’s methods and as outlined previously (24). Suitable riboprobes (sense and antisense probes) were prepared by PCR (supplemental Table S1), cloned, and transcribed using T7 RNA polymerase. To label the RNA probes, digoxigenin-11-UTP (Roche) was incorporated using the AmpliScribe T7 kit (Epicenter Technologies). Treated embryos and corresponding controls were fixed in 4% p-formaldehyde, dehydrated in graded ethanol/xylene mixtures and then embedded in paraffin (24). Sagittal sections (5 μm thick) of the embryos were prepared and stored at room temperature prior to in situ hybridization.

Immunohistochemistry—Immunohistochemical staining was performed on paraffin-embedded embryo sections employing the immunoperoxidase method. The antigenic epitopes were first exposed by heating the sections in 10 mM citrate buffer (pH 6.0) containing 0.05% Tween 20. After quenching the endogenous peroxidase with 3% hydrogen peroxide, sections were incubated in TBS (5% normal sheep serum) containing either anti-human MAO-A (clone C-19) or anti-human MAO-B (clone C-17) monoclonal antibodies (Santa Cruz Biotechnology, Inc.). The sections were then incubated with a peroxidase-labeled secondary antibody (Biocare Medical Kit), and slides were developed with diaminobenzidin (brown color). Negative controls lacked the primary antibodies.

Apoptotic cells were stained with both the supravital dye staining method and standard TUNEL technique. The whole mount supravital dye Nile blue sulfate visualizes non-viable cells. After cultivation, embryos at E10.5 were washed in Ringer’s solution, bathed in a Nile blue sulfate solution for 15–20 min at 37 °C, and then washed in Ringer’s solution before microscopic examination. TUNEL staining was performed on embryo sagittal sections using the Chemicon kit according to the vendor’s instructions. Nuclear DNA was counterstained with 0.5% (w/v) methyl green for microscopic examination.

Western Blotting—For caspase 3, caspase 8, caspase 9, cyclin D1, E2F1, and β-actin immunoblotting, whole embryo lysates were obtained by homogenizing embryos in lysis buffer (50 mM Tris HCl, 0.3 M NaCl, 1 mM EDTA, 1 mM DTT, and 1 mM PMSF). Aliquots with equal amounts of total protein (100 μg) were separated on 12% SDS-PAGE gels, and separated proteins were transferred to nitrocellulose membranes (Amersham Biosciences). The membranes were blocked overnight with an ovalbumin solution (Sigma) and then incubated at 37 °C for 1 h with primary antibodies. The membranes were then washed three times with TBS and incubated with HRP-labeled secondary antibody (Dako) at 1:1000 dilution (30 min at room temperature). Blots were finally developed using the ECL kit (GE Healthcare). Immunoreactive protein bands were imaged using the Quantity One® quantification software (Bio-Rad) and normalized to β-actin expression. For MAO and R1 immunoblotting, primary antibodies were purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA), but for quantification of caspase 3, caspase 8, caspase 9, and β-actin, antibodies were obtained from Abcam, Inc. (Cambridge, UK).

Serotonin Assay—Serotonin concentration in the embryos was measured using the Serotonin enzyme immunoassay (Ultra Sensitive) kit (Immuno Biological Laboratory, Inc.) according to the manufacturer’s instructions. In brief, after treatments in vitro cultured embryos were washed and homogenized, and aliquots of the lysates were applied for quantification. The lower detection limit of the kit was 1.0 nmol/liter, and a linear calibration plot was established between 1.45 and 100 nmol/liter ($r^2 = 0.9807 \pm 0.0012$). The intra- and interassay variation coefficients were < 3.02%, and the recovery range varied between 100.4 and 102.3%.

RESULTS

Expression of MAO mRNA During the Time Course of Murine Embryogenesis—Previous in situ hybridization indicated that MAO isoforms are expressed in embryonic mouse brain as early as at E12 (17). Employing a more sensitive quantitative RT-PCR (RT-qPCR) strategy, we detected MAO mRNA species much earlier on during mouse embryogenesis. At E6.5, MAO-A mRNA was clearly detectable, and its concentration increased until E9.5. Then it remained at constant levels until E13.5 (Fig. 1A). At E14.5, MAO-A mRNA levels dropped and did not recover until birth. MAO-B mRNA follows similar expression kinetics, but its average steady state concentration was 4-fold lower than MAO-A mRNA. The prevalence of MAO-A over MAO-B in murine embryos was confirmed by activity assays (Fig. 1B). More than 90% inhibition of total MAO activity was observed when total embryonic lysates were assayed in the presence of 10^{-7} M clorgyline, a specific MAO-A inhibitor. In contrast, when the MAO-B-specific inhibitor deprenyl was used, a similar degree of inhibition was only observed at 10^{-5} M.

To follow cerebral expression of MAO isoforms during embryo development, we prepared embryonic brains at different developmental stages and recorded the concentration profiles for MAO mRNAs (Fig. 1C). Cerebral MAO-A mRNA follows similar expression kinetics as in the whole embryo, with constant levels from E10.5 to E13.5, and dropped down to lower levels when the embryos approached birth. MAO-B mRNA concentrations are lower than those of MAO-A and decline further toward delivery.

To explore the spatial distribution of MAO mRNA expression during embryogenesis, we performed in situ hybridization (Fig. 2). Between E10.5 and E12.5, specific signals for MAO-A mRNA expression (blue staining indicated by the red arrows) were mainly observed in developing brains. Low levels of in situ hybridization signals for MAO-B mRNA were detected in the fore- and hindbrain at E10.5. At later stages (E11.5 and E12.5), the signals were even less intense.

Silencing of MAO-A Expression Induced Developmental Defects—To silence expression of MAO-A and MAO-B during in vitro embryogenesis, we designed different specific siRNA probes for each MAO-isoform (supplemental Table S1). In an initial experiment, different siRNA species were injected into the amniotic cavity, and embryos were recovered after a cultur-
ing period of 72 h. Following knockdown of MAO-A using two independent siRNA constructs, we observed severe developmental retardations for both siRNA species, as indicated by the reduced embryo size compared with embryos treated with scrambled siRNAs (Fig. 3A). In contrast, embryos treated with siRNAs targeted at MAO-B expression did not have an apparent impact on embryo development (data not shown).

To characterize the siRNA-induced developmental defects in different organ systems in more detail, we employed a morphometric scoring system (25). This scoring system enabled us to translate the developmental progress at organ and tissue levels into a quantifiable matrix of scoring points. Using this system, we evaluated the siRNA-induced developmental defects of the developing brain, heart, and limbs in addition to general growth retardations (crown rump length). We found that siRNA-mediated knockdown of MAO-A expression reduced
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FIGURE 4. Impact of MAO expression silencing on embryo development. For knockdown studies, siRNA constructs or corresponding control probes were mixed with an appropriate volume of transfection reagent (see "Materials and Methods"). Murine embryos were explanted at E7.5, and the transfection mixture (10 nl with 50 nm probe) was injected into the amniotic cavity. The embryos were then maintained in in vitro cultures for 2 h (E7.5), 24 h (E8.5), 48 h (E9.5) and 72 h (E10.5), and silencing of MAO expression was tested. Whole mount in situ hybridization: Mouse embryos were removed from in vitro culture and in situ hybridization with MAO mRNA antisense probes was carried out. The red arrows indicate MAO mRNA expression. The blue arrows indicate developmental defects of the brain. fb, forebrain; mb, midbrain; hb, hindbrain; hf, headfold; tb, tail bud.

The success of the knockdown strategy was verified by whole mount in situ hybridization. Strong expression of both MAO isoforms was observed in the head region, as indicated by the dark blue in situ hybridization signals (Fig. 4). Embryos treated with siRNA species targeted at MAO-A expression exhibited strongly reduced in situ hybridization signals after 72 h incubation (Fig. 4, panels 1–4). Similar results were observed when embryos were treated with MAO-B siRNA (Fig. 4, panels 5–8). These data suggest successful knockdown of the expression of either MAO isoform.

Next, we aimed to analyze the effects of MAO knockdown at different time points during our experimental time frame. Both MAO isoforms are expressed at E7.5 and E8.5 in the headfold region and in the caudal neural tube, as indicated by the dark blue in situ hybridization signals (Fig. 4). At these developmental stages injection of siRNA constructs strongly reduced expression of the corresponding MAO isoform without causing apparent developmental alterations. At E9.5 mRNA of both MAO isoforms was mainly detected in the developing fore-, mid-, and hindbrain and the forelimbs. Faint in situ hybridization signals found in the developing heart indicate expression of MAO-A in this organ (Fig. 4, red arrows). Treatment with specific siRNA selectively reduced MAO isoform expression. However, at this stage (Fig. 4, upper panel), we observed developmental defects as indicated by the reduced size of the whole embryo and of the developing brain (blue arrow). In conclusion, developmental retardations induced by MAO-A siRNA are apparent by day E9.5 but become most pronounced at day E10.5.

To confirm siRNA-mediated knockdown of MAO isoforms on the protein level, we performed immunohistochemistry and Western blotting (Fig. 5, A and B). Here we detected isoform-specific and dose-dependent siRNA-induced reduction of MAO protein. This indicates successful down-regulation of target gene expression as well as a high isoform specificity of our siRNA constructs.

To make sure that our knockdown data is not the results of methodological artifacts (off-target effects of the siRNA constructs) we employed two additional experimental approaches. Firstly, we used the specific and irreversible inhibitor of MAO-A activity clorgyline. Secondly, we transfected the developing embryos with a mammalian expression plasmid that drives the overexpression of the transcriptional inhibitor of MAO-A expression R1. Indeed, our Western blotting data indicates specific suppression of MAO-A but not of MAO-B expression following R1 overexpression (Fig. 5C).

Reduced expression and/or activity of MAO-A has been reported to be associated with increased systemic levels of its substrate serotonin (13, 15). This prompted us to measure embryonic serotonin levels following MAO-A knockdown. Indeed, we observed that embryonic serotonin levels were elevated almost 4-fold 72 h after MAO-A siRNA treatment (Fig. 5D). Similarly, serotonin levels were also elevated when MAO-A activity was inhibited by clorgyline or when MAO-A gene transcription was suppressed by R1 overexpression.

Inhibition of MAO-A Activity Induces Developmental Defects Similar to MAO-A Knockdown—If the developmental abnormalities observed following siRNA-mediated knockdown of MAO-A expression are specific, pharmacological inhibition of MAO-A activity or suppression of MAO-A gene expression by R1 overexpression should induce similar developmental alterations as siRNA treatment. Indeed, when embryos were cultured in the presence of clorgyline or when R1 was overexpressed, embryo growth was impaired to a similar extent as observed for MAO-A knockdown embryos (Fig. 6, A and C). Clorgyline treatment resulted in retarded development of the hind- and midbrain, whereas development of the forebrain was less affected (Fig. 6B). Development of the limbs and the heart proceeded normally. Similar observations were made when R1
was overexpressed (Fig. 6D). Thus, embryos treated with clorgyline or in whom R1 was overexpressed phenotypically resemble MAO-A knockdown embryos.

Knockdown of MAO-A Expression Inhibited Developmental Apoptosis—Apoptosis is a vital process in embryo development (27) and MAO-A has been implicated in the regulation of cell proliferation and programmed cell death (26, 28). To test whether knockdown of MAO-A expression impacts developmental apoptosis during murine embryogenesis, we first monitored embryonic cell death by whole mount Nile blue staining of embryos at E10.5 (30-somite stage) and by the TUNEL technique in microscopic cross-sections (Fig. 7A). When we knocked down embryonic expression of MAO-A, the number of dead (Nile blue staining) and apoptotic cells (TUNEL staining) in the neuroepithelium of the developing brain was decreased significantly (yellow and red arrows, respectively) (Fig. 7A).

For more detailed information on the mechanisms of the developmental defects, we repeated the knockdown experiments and evaluated the impact of the siRNA constructs on the activation state of various caspases (Fig. 7, B and C). MAO-A is thought to modulate the intrinsic branch of the apoptotic pathway involving caspase 3 and caspase 9 (26). When we knocked down embryonic MAO-A expression, the activation states of caspase 3 and caspase 9 were reduced significantly, whereas activation of caspase 8 was not altered (Fig. 7, B and C). These
data suggest that knockdown of MAO-A expression reduces apoptotic cell death by interfering with the intrinsic pathway of apoptosis.

Neural development relies on the fine balance between apoptotic cell death on one hand and cell proliferation and differentiation on the other (29). MAO-A has been shown to modulate cell proliferation by regulating intracellular levels of cyclin D1 and E2F1 (28). Cyclin D1 specifically associates with selected cyclin-dependent kinases, phosphorylating the retinoblastoma protein 1 (Rb1). This phosphorylation suppresses the growth inhibitory activity of hypophosphorylated Rb1 and thus induces cell proliferation by releasing E2F1 (30). We found that MAO-A knockdown was paralleled by a reduction in cellular cyclin D1 levels (Fig. 7B). In contrast, cellular levels of the transcriptional regulator E2F1 whose activity is associated with proliferation (31) remained unaffected by any treatment (Fig. 7B). Similar observations were made when MAO-A activity was inhibited using clorgyline or when MAO-A expression was suppressed by R1 overexpression.

**DISCUSSION**

In vertebrates MAO isoforms metabolize neurotransmitters and thus are important for the functionality of the adult brain. However, the role of MAO in murine embryonic development is less well understood. To fill this gap of knowledge, we employed an in vitro embryogenesis model (24). Mouse strains that exhibit incidental inactivation of the MAO-A gene are fertile and only show minor developmental alterations (13, 14), which appears to conflict with the robust changes we observed. However, our experimental setup avoids limitations of stem cell-based approaches, such as adaptive or compensational effects (23). Such effects may be responsible for the lack of developmental defects in MAO-A-deficient mouse strains (13, 14). Moreover, our experimental approach causes a sudden reduction of MAO-A expression at a certain developmental stage in embryos that have grown in the presence of MAO-A until expression knockdown was initiated. Knockout embryos, in contrast, might have activated compensatory mechanisms during early embryo development that are not available anymore at later developmental stages. In this respect, the two experimental models (knockout versus knockdown) are different and may lead to different experimental data. To make sure that our knockdown data are not the result of methodological artifacts, we combined genetic (siRNA-mediated MAO-A knockdown and R1 overexpression) and pharmacological (MAO-A inhibitor clorgyline) approaches and obtained identi-
cal results. In fact, recent observations of impaired cell proliferation of the telencephalon during late embryo development in an MAO-A/B-deficient mouse line supports our line of argumentation that MAO-A affects prenatal development (32).

Our studies indicate that both MAO-A and MAO-B are expressed at early stages when neuronal differentiation (gastrulation and neurulation) is induced and major subdivision of the brain (migration, specification, segmentation) takes place (33). MAO-A is a proapoptotic enzyme, and caspase 3 and caspase 9 of the intrinsic pathway of apoptosis have been related to MAO-A activity (26, 34). Moreover, MAO-A appears to modulate cell proliferation via controlling cytoplasmic cyclin D1 and E2F1 levels (28). The precise mechanisms of MAO-mediated regulation of cellular signaling cascades have not been explored in detail. A potential effector molecule is hydrogen peroxide, which is generated by MAO activity and affects both mitochondrial function and the cellular redox state (35, 36). On one hand, mitochondrial dysfunction is a feature of the intrinsic pathway of cell death signaling and is paralleled by increased lipid peroxidation of the mitochondrial membranes (37). On the other hand, changes of the cellular redox state affect a multitude of signaling cascades involved in the regulation of cellular proliferation and differentiation (38). However, more research is needed to understand the role of MAO-A derived hydrogen peroxide in the regulation of these signaling events.

Moreover, MAO-A has been implicated in cell cycle progression mediated by the oncogene c-myc (28). The myc protein acts as a transcription factor that modulates the expression of a variety of genes associated with cell cycle regulation, including E2F1 and cyclin D1 (39–41). In addition, c-myc regulates the expression as well as the activity of the transcription factor R1 (28). We observed similar developmental abnormalities in embryos overexpressing R1 compared with MAO-A knockdown embryos. Moreover, R1 overexpression similar to MAO-A knockdown or pharmacological inhibition of MAO-A activity coincides with elevated levels of the cyclin D1 protein. In conclusion, this data implicates MAO-A as an integral mediator of the c-myc proliferation signaling cascade via modulating cyclin D1 levels.

Developmental apoptosis is a well balanced process that is crucial for the formation of embryonic structures. Disturbance of this equilibrium induces morphological abnormalities, such as neural tube defects (29, 42). We found that MAO-A knockdown embryos and embryos exposed to clorgyline showed normal neural tube closure, which was followed by decreased apoptosis as well as impaired cell proliferation. Embryos of caspase
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3-deficient mice die before birth and exhibit profoundly impaired brain development and cellular hyperplasia (43). In our MAO-A knockdown studies, caspase 3 activation was significantly impaired, which is in line with caspase 3 knockout data.

A hallmark of MAO-A deficiency in previously reported animal models are elevated levels of serotonin (13). We also observed elevated embryonic serotonin levels following MAO-A knockdown or following pharmacologic inhibition of MAO-A activity. These data underline the validity of the different experimental setups. Elevated 5-HT levels were also found in neural stem cells treated with Moclobemide, a reversible inhibitor of MAO-A activity. MAO-A inhibition resulted in an induction of the expression of the anti-apoptotic factor Bcl-2 in neural stem cells via ERK activation, and thereby inhibited apoptotic signaling and subsequently induced stem cell differentiation (34). Newborn mice lacking the cerebral vesicular monoamine transporter (VMAT2<sup>−/−</sup>) suffer from increased developmental apoptosis in the cerebral cortex (44). In these animals, caspase 3 and caspase 9 are hyperactive, and expression of antiapoptotic Bcl-2 was reduced. When these mice were crossed with MAO-A deficient animals (VMAT2<sup>−/−</sup>/MAO-A<sup>−/−</sup> double KO mice), 5-HT levels were elevated, and the increased cell death induced by VMAT2 deficiency was normalized (44). This finding is consistent with our data and also implicates MAO-A in proper brain development.

Although our data suggest MAO-A as an important player in the regulation of cerebral embryogenesis, more work is needed to identify the components of the associated signaling cascades. For instance, dysregulation of the finely tuned spatial and temporal balance between 5-HT production and breakdown may have fatal consequences for the developing embryo. Indeed, for several years it has been a matter of discussion of whether or not pharmacological intervention aimed at 5-HT homeostasis, such as selective serotonin reuptake inhibitors, during pregnancy has teratogenic effects (45, 46).

REFERENCES

1. Shih, J. C. (2007) Neurochem. Res. 32, 1757–1761
2. Scrutton, N. S. (2004) Nat. Prod. Rep. 21, 722–730
3. Bach, A. W., Lan, N. C., Johnson, D. L., Abell, C. W., Bembenek, M. E., Kwan, S. W., Seeburg, P. H., and Shih, J. C. (1988) Proc. Natl. Acad. Sci. U.S.A. 85, 4934–4938
4. Gheva, R. M., Rebrin, L., Chen, K., and Shih, J. C. (2001) J. Biol. Chem. 276, 9877–9882
5. Jay, J. W., Houpt, T. A., Wessel, T. C., Chen, K., Shih, J. C., and Joh, T. H. (1997) Synapse 25, 30–36
6. Billett, E. E. (2004) Neurotoxicology 25, 139–148
7. Chen, K. (2004) Neurotoxicology 25, 31–36
8. Chen, K., Ou, X. M., Chen, G., Choi, S. H., and Shih, J. C. (2005) J. Biol. Chem. 280, 11592–11599
9. Syagailo, Y. V., Stöber, G., Grässel, M., Reimer, E., Knapp, M., Jungkunz, G., Okladnova, O., Meyer, J., and Lesch, K. P. (2001) Am. J. Med. Genet 105, 168–171
10. Suarez-Merino, B., Bye, J., McDowall, J., Ross, M., and Craig, I. W. (2001) Hum. Mutat. 17, 523
11. Youdim, M. B., Edmondson, D., and Tipton, K. F. (2006) Nat. Rev. Neurosci. 7, 295–309
12. Grimsby, J., Toth, M., Chen, K., Kumazawa, T., Klaidman, L., Adams, J. D., Karoum, F., Gal, J., and Shih, J. C. (1997) Nat. Genet. 17, 206–210
13. Cases, O., Seif, I., Grimsby, J., Gaspar, P., Chen, K., Pourrain, S., Müller, U., Aguet, M., Babinet, C., and Shih, J. C. (1995) Science 268, 833–875
14. Scott, A. L., Bortolato, M., Chen, K., and Shih, J. C. (2008) Neuroreport 19, 739–743
15. Chen, K., Holschneider, D. P., Wu, W., Rebrin, I., and Shih, J. C. (2004) J. Biol. Chem. 279, 39645–39652
16. Luo, X., Persico, A. M., and Lauder, J. M. (2003) Dev. Neurosci. 25, 173–183
17. Vitalis, T., Fouquet, C., Alvarez, C., Seif, I., Price, D., Gaspar, P., and Cases, O. (2002) J. Comp. Neurol. 442, 331–347
18. Burnet, H., Bevengut, M., Chakri, F., Bou-Flores, C., Coulon, P., Gaytan, S., Pasaro, R., and Hilaire, G. (2001) J. Neurosci. 21, 5212–5221
19. Upton, A. L., Salichon, N., Lebrand, C., Ravary, A., Blakely, R., Seif, I., and Gaspar, P. (1999) J. Neurosci. 19, 7007–7024
20. Popova, N. K., Maslova, L. N., Morosova, E. A., Bulygina, V. V., and Seif, I. (2006) Psychoneuroendocrinology 31, 179–186
21. Chen, K., Cases, O., Rebrin, I., Wu, W., Gallaher, T. K., Seif, I., and Shih, J. C. (2007) J. Biol. Chem. 282, 115–123
22. Beglopooulos, V., and Shen, J. (2004) Neuromolecular Med. 6, 13–30
23. Senechal, Y., Larmet, Y., and Dev, K. K. (2006) Neurodegener Dis. 3, 134–147
24. Ufer, C., Wang, C. C., Fähling, M., Schiebel, H., Thiele, B. I., Billett, E. E., Kuhn, H., and Borchert, A. (2008) Genes Dev. 22, 1838–1850
25. Van Maele-Fabry, G., Delhaese, F., and Picard, J. J. (1990) Toxicol. In Vitro 4, 149–156
26. Fitzgerald, J. C., Ufer, C., De Girolamo, L. A., Kuhn, H., and Billett, E. E. (2007) J. Neurochem. 103, 2189–2199
27. Bortner, C. D., and Cidlowski, J. A. (2002) Annu. Rev. Pharmacol. Toxicol. 42, 259–281
28. Ou, X. M., Chen, K., and Shih, J. C. (2006) Proc. Natl. Acad. Sci. U.S.A. 103, 10923–10928
29. Copp, A. J. (2005) J. Anat. 207, 623–635
30. Guo, Y., Yang, K., Harwalkar, J., Nye, J. M., Mason, D. R., Garrett, M. D., Hitomi, M., and Stacey, D. W. (2005) Oncogene 24, 2599–2612
31. McClellan, K. A., and Slack, R. S. (2007) Cell Cycle 6, 2917–2927
32. Cheng, A., Scott, A. L., Ladenheim, B., Chen, K., Ouyang, X., Lathia, J. D., Mughal, M., Cadet, J. L., Mattson, M. P., and Shih, J. C. (2010) J. Neurosci. 30, 10752–10762
33. Wang, C. C. (2005) Neuroembryol. Aging 3, 78–91
34. Chouiti, S. H., Hu, H. T., Lin, H. L., Chen, L. H., Chien, C. S., Ho, L. L., Lee, C. H., and Chang, Y. L. (2006) Br. J. Pharmacol. 148, 578–597
35. Naoi, M., Maruyama, Y., Hi, H., Inaba, K., Akao, Y., and Shimato-Nagai, M. (2009) J. Neural Transm. 116, 1371–1381
36. Cohen, G., Farooqui, R., and Kesler, N. (1997) Proc. Natl. Acad. Sci. U.S.A. 94, 4890–4894
37. Imai, H., and Nakagawa, Y. (2003) Free Radic. Biol. Med. 34, 145–169
38. Ufer, C., Wang, C. C., Borchert, A., Heydeck, D., and Huhn, K. (2010) Antioxid. Redox Signal. 14, 2917–2927
39. Kuida, K., Yamasaki, Y., and Kusumi, K. (2004) Cell 116, 1371–1381
40. Santoni-Rugiu, E., Jensen, M. R., and Thorgeirsson, S. S. (1998) Cancer Res. 58, 123–134
41. Seif, I., Grimsby, J., and Shih, J. C. (2008) Cell Death Differ. 15, 1170–1177
42. Stankovski, L., Alvarez, C., Ouimet, T., Vitalis, T., El-Hachimi, K. H., Price, D., Deneris, E., and Flavell, R. A. (1996) Nature 384, 368–372
43. Alwan, S., and Friedman, J. M. (2009) CNS Drug 23, 493–509
44. Borue, X., Chen, J., and Condron, B. G. (2007) Int. J. Dev. Neurosci. 25, 341–347