RESEARCH ARTICLE

Zebrafish duox mutations provide a model for human congenital hypothyroidism

Kunal Chopra, Shoko Ishibashi and Enrique Amaya*

ABSTRACT

Thyroid dyshormonogenesis is a leading cause of congenital hypothyroidism, a highly prevalent but treatable condition. Thyroid hormone (TH) synthesis is dependent on the formation of reactive oxygen species (ROS). In humans, the primary sources for ROS production during thyroid hormone synthesis are the NADPH oxidases DUOX1 and DUOX2. Indeed, mutations in DUOX1 and DUOX2 have been linked with congenital hypothyroidism. Unlike humans, zebrafish has a single orthologue for DUOX1 and DUOX2. In this study, we investigated the phenotypes associated with two nonsense mutant alleles, sa9892 and sa13017, of the single duox gene in zebrafish. Both alleles gave rise to readily observable phenotypes reminiscent of congenital hypothyroidism, from the larval stages through to adulthood. By using various methods to examine external and internal phenotypes, we discovered a strong correlation between TH synthesis and duox function, beginning from an early larval stage, when T4 levels are already noticeably absent in the mutants. Loss of T4 production resulted in growth retardation, pigmentation defects, ragged fins, thyroid hyperplasia/external goiter and infertility. Remarkably, all of these defects associated with chronic congenital hypothyroidism could be rescued with T4 treatment, even when initiated when the fish had already reached adulthood. Our work suggests that these zebrafish duox mutants may provide a powerful model to understand the aetiology of untreated and treated congenital hypothyroidism even in advanced stages of development.

This article has an associated First Person interview with the first author of the paper.

KEY WORDS: Congenital hypothyroidism, Growth retardation, Infertility, Thyroid

INTRODUCTION

Congenital hypothyroidism (CH) is an endocrine disorder that may result from disrupted thyroid hormone (TH) synthesis (15–20% of all cases) or impaired development of the thyroid gland [thyroid dysgenesis (TD)] (80% of all cases) (Kizys et al., 2017). CH is the most prevalent congenital endocrine disorder and is believed to be one of the most preventable causes of mental retardation (Chakera et al., 2012; Olivieri, 2015; Roberts and Ladenson, 2004). Indeed, in infants younger than 3 months of age, neurological damage progressively worsens with delay in starting treatment (Virtanen et al., 1983). Mutations in the NADPH oxidase DUOX2 and, to a lesser extent, DUOX1 have been associated with dyshormonogenesis in CH patients (Aycan et al., 2017; Moreno et al., 2002). DUOX1 and DUOX2 generate hydrogen peroxide (H2O2), which is a crucial electron acceptor during thyroid peroxidase-catalysed iodiumination and coupling reactions occurring while TH synthesis is underway (De Deken et al., 2000; Dupuy et al., 1999). H2O2 production is a limiting step in TH biosynthesis. The main source of H2O2 in the thyroid is DUOX2 in conjunction with its maturation factor DUOX2A, both of which are located at the apical surface of the thyroid follicular cells, thyrocytes. DUOX2-mediated H2O2 acts as a thyreroxidase (TPO) co-substrate, rapidly oxidising iodine and resulting in its covalent binding to the tyrosine residues of thyroglobulin in the follicular lumen. This produces moniodotyrosine (MIT) and diiodotyrosine (DIT), in the thyroglobulin molecule, which undergo coupling to give the THs triiodothyronine (T3) and thyroxine (T4) (Carvalho and Dupuy, 2013; Muzza and Fugazzola, 2017; Sugawara, 2014). A negative feedback loop is in charge of thyroid size and function. Thyrocytes secrete T3 and T4 and these inhibit the production of the thyroid-stimulating hormone (TSH) via the anterior pituitary thyrotropes (Dumont et al., 1992). Thyrocytes respond to limiting physiological stimuli by a way of hypertrophy and proliferation. This is a direct response to compensate for diminishing THs in conditions including, but not limited to, iodine deficiency, exposure to anti-thyroid drugs and punctuated production of reactive oxygen species (ROS). It has been shown that early initiation of TH treatment (within 3 weeks post-partum) leads to normal IQ and physical growth and correlates with excellent prognoses (Aronson et al., 1990; Clause, 2013; Rahmani et al., 2016; Rovet et al., 1987). Expectedly then, if treatment is delayed beyond 4 weeks, individuals become increasingly prone to mental retardation and incomplete physical growth (Gilbert et al., 2012; Zimmermann, 2011). To date, various approaches have been adopted to induce hypothyroidism in animal models, including surgical removal of the thyroid gland, thyroid gland removal via radioactive iodine isotope (131I), dietary restriction of iodine, and goitrogen administration (Argumedo et al., 2012). We present here a zebrafish model of CH, which exhibits several phenotypes associated with CH in humans, including growth retardation. Interestingly, while CH zebrafish display growth retardation initially, they are able to reach normal size eventually without the need for pharmacological intervention. The additional external and internal phenotypes associated with hypothyroidism are restored upon treatment with T4, including restoration of reproductive function, even when treatment is applied during adulthood.

RESULTS

Molecular characterisation of duox mutant alleles

Duox is a member of the NADPH oxidase (NOX) family of enzymes. Seven NOX family members are present in the human...
amino acid protein. In order to investigate the function of zebrafish C-terminal NADPH and FAD binding sites, they would be expected to examine truncations of the Duox protein, including the loss of the two critical amino acids. Since both these premature codons result in a C>T change (Fig. 1C) creating a premature stop codon (TGA) after the 944th amino acid; the second allele, sa13017, contains a nonsense mutation in exon 23, resulting in a C>T change (Fig. 1B) and a premature stop codon (TAG) after the 944th amino acid; the second allele, duox sa13017, contains a nonsense mutation in exon 23, resulting in a C>T change (Fig. 1C) creating a premature stop codon (TGA) after the 997th amino acid. Since both these premature codons result in truncations of the Duox protein, including the loss of the two critical C-terminal NADPH and FAD binding sites, they would be expected to be loss-of-function mutations. Genotyping of these alleles can be performed via genomic PCR followed by sequencing of these regions (Fig. 1B,C).

**duox mutants are growth retarded**

We in-crossed duox sa9892+/− and duox sa13017+/− sibling adults and inter-crossed duox sa9892+/− with duox sa13017+/− adults to produce a range of wild-type (WT), heterozygous, homozygous mutant and compound heterozygous mutant animals containing both alleles. While the WT, duox sa9892+/− and duox sa13017+/− animals were phenotypically indistinguishable, the homozygous mutants of both alleles, and the compound mutants (i.e. duox sa9892/sa13017) displayed a number of phenotypes that were distinct from those seen in the WT and heterozygous siblings. The first overtly apparent phenotype exhibited by the duox sa9892+/−, the duox sa13017+/− and the trans-heterozygous duox sa9892/sa13017 mutants was that they were growth retarded. At 3 months of age, the sa9892+/−, sa13017+/− and sa9892/sa13017 mutant fish were significantly shorter in terms of body length than their WT and heterozygous siblings (Fig. 2A–G). At 6 months of age, the duox sa9892+/− and duox sa13017/sa13017 mutant animals caught up in size with their WT and heterozygous siblings. However, the duox sa13017+/− animals still remained stunted (Fig. 2H). Another phenotype suggestive of slowed growth was apparent in the growth and organogenesis of the swim bladder. The swim bladder is a hydrostatic organ, which becomes bi-lobed by 21 dpf (Winata et al., 2009). We found that the swim bladder in the duox sa9892−/− animals remained unilobed even at 54 dpf (n=9) (Fig. 2I,J). Homozygous duox mutants also exhibit a delay or absence of development of barbels, which are a set of anterior sensory appendages. Zebrafish develop a short pair of nasal barbels and a long trailing pair of maxillary barbels (Fig. 2K,N). These are normally visible by 1 month post-fertilisation and sustained throughout life (LeClair and Topczewski, 2010). In all cases, homozygous duox mutants lacked barbels at 3 months of age (Fig. 2L,O,P). However, between 6 and 10 months of age, maxillary barbels were seen in some older duox sa9892−/− (five out of 11; see Fig. 2M) and sa9892/sa13017 (two out of 11) animals, but in none of the sa13017−/− animals (zero out of nine).

**duox mutants have dark pigmentation, erythema and ragged fins**

Zebrafish are recognisable by their eponymous pattern of five dark blue stripes alternating with four lighter yellow inter-stripes, covering the lateral flanks, and anal and caudal fins (Singh and Nüsslein-Volhard, 2015). The stripes are comprised of black melanophores with a few iridescent iridophores, while the inter-stripes are comprised of yellow and orange xanthophores and numerous iridophores (Hirata et al., 2003). We found that the homozygous duox mutants were darker than their WT and heterozygote siblings (Fig. 3A–C). The darker pigmentation was associated with the presence of approximately twice the number of melanophores in the homozygous duox mutants, relative to their heterozygous and WT siblings (Fig. 3D). Conveniently, we also found that the difference in pigmentation was sufficient to allow for phenotypic identification of homozygous duox mutants from their heterozygous and WT siblings, with 100% accuracy, as confirmed retrospectively via genotyping. In addition, homozygous duox mutants also showed stripe irregularities not seen in WT and heterozygous siblings, such as wavy stripes and stripe discontinuities (Fig. 3E,F). Thus, pigmentation differences can be used as a reliable identification method for distinguishing homozygous duox mutants from their heterozygous and WT siblings, as early as 60 dpf.

Less apparent but nevertheless significant were craniofacial anomalies among adult mutants. In particular, we found a significant shortening of the frontal height among the duox sa9892−/−, sa13017−/− and sa9892/sa13017 animals, when compared to their WT and heterozygous siblings (Fig. 3G–I). Adding further to the list of phenotypes, we noticed erythema (redness) in the opercular region of mutants (Fig. 3J–O). This was especially prominent in background strains that lack melanophores, such as nacre and casper. The redness was most apparent in juvenile fish.

Finally, the homozygous duox mutants often displayed misshapen or damaged fins (Fig. 3P–U). We found that the duox sa9892−/− (15 out of 15), sa13017−/− (seven out of seven) and sa9892/sa13017 (19 out of 23) animals displayed damaged fins. In many cases this was manifested as vertical (dorsal and anal fin) or horizontal (caudal fin) tears in the fins. In other cases, there were spontaneous losses of portions of fins or ragged fin margins (Fig. 3S–U). Damaged fins were noticeable as early as 42 dpf.

**Homozygous duox mutants are viable but are unable to breed**

While we found that duox sa9892−/− and sa13017−/− mutants reached adulthood, unlike their heterozygous and WT siblings, they were unable to breed. Females, although gravid, were found not to lay eggs regardless of pairing with mutant, heterozygous or WT males. Similarly, mutant males failed to cross with females, regardless of genotype. Furthermore, we noticed that homozygous duox mutant females seemed egg-bound, suggesting that they were unable to lay eggs (Fig. 4A,B; left panels). We confirmed that females do contain eggs internally via histological sectioning (Fig. 4A,B; right panels) as well as by abdominal squeezing to release eggs. Similarly, compound heterozygotes of the two alleles were found to be viable but failed to breed, and the females also became egg bound.
Homozygous duox mutants develop goitres

In addition to the phenotypes described above, we noted that some of the homozygous duox mutant adult animals displayed external goitre-like growths in the submandibular area (Fig. 5A,B). These external goitre-like growths were observed among adults older than 3 months of age, across all three mutant combinations – duox sa9892+/− (11 out of 18), sa13017+/− (six out of 11) and sa9892/sa13017 (three out of 22). These richly vascularised growths were variously sized. Additionally, some of the animals also exhibited lateral flaring of opercular flaps (Fig. 5C). To confirm whether these goitre-like growths were indeed enlarged thyroids, we fixed and sectioned a subset of growths were indeed thyroid origin (Fig. 5D–L). Also, the thyroid hyperplasia was in striking contrast to the size of the thyroids in the WT and heterozygous mutant siblings, and performed in situ hybridization (ISH) analysis for the expression of thyroglobulin, a thyroid marker. The results confirmed that these growths were indeed of thyroid origin (Fig. 5D–L). Also, the thyroid hyperplasia was in striking contrast to the size of the thyroids in the WT and heterozygous mutant siblings, where the extent of thyroglobulin staining was much smaller and more distinct, presenting as discreet rings confined to the ventral mid-pharyngeal region (Fig. 5E,F). Importantly, all mutants had internal thyroid hyperplasias, regardless of the presence or absence of external goiters (Fig. 5G–L).

duox mutants are in a state of hypothyroidism

The goitre-like growths, as well as the other phenotypes observed in the homozygous duox mutants suggested that the mutants might be exhibiting hypothyroidism. To test whether this might be the case, we assessed the presence of thyroxine (T4) in the homozygous duox mutants and in their heterozygous and WT siblings via wholemount immunostaining. We found that, while the WT and heterozygous siblings exhibited robust T4 staining (Fig. 6A–C), duox sa9892+/− and sa13017+/− larvae had no detectable T4 staining in their thyroids (Fig. 6D,E). Consistent with the loss of T4 being due to lack of NADPH oxidase activity in the homozygous duox mutants, we were able to phenocopy the loss of T4 staining in the larvae by treating them with the NADPH oxidase inhibitor, diphenyleneiodonium (DPI) (Fig. 6F).

duox-mediated hypothyroidism is responsive to T4 treatment

Among humans, CH responds very well to T4 treatment, especially when treatment is initiated as soon as hypothyroidism is suspected (Rahmani et al., 2016). Here, we decided to ask whether supplementation of the aquarium water with T4 could reverse some or all of the phenotypes observed in the homozygous duox mutants. We initiated T4 (30 nM) treatment of the duox sa9892+/− and sa9892+/− animals starting at 11 months of age, when all of the phenotypes described previously were already apparent. We found that most of the phenotypes associated with loss of duox function could be reversed by treatment with T4. Body pigmentation was the first phenotype to be reversed in the treated animals, such that by 2 weeks after the initiation of treatment the duox sa9892+/− animals became visibly paler than their untreated duox sa9892+/− siblings (Fig. 7A versus 7B). The difference in pigmentation was associated with a significant decrease in melanophore density in T4-treated homozygous mutant animals when compared to the untreated homozygous mutant animals (Fig. 7C–G). Indeed, the density of melanophores in the treated mutants was similar to that seen in untreated or treated heterozygous mutant animals, suggesting a complete rescue (Fig. 7C–G). In addition, we found that fin quality improved markedly, with treated mutants showing fuller, unbroken fins compared to the ragged fins of the untreated controls (compare Fig. 7A and B). Furthermore, after 8 weeks of T4 treatment we were able to rescue breeding behaviour in both sexes. Mutant males and females were able to spawn with WT animals or with each other. These mating episodes resulted in the production of four clutches of eggs in four consecutive weeks. Rescue of fertility was perhaps the most striking outcome of T4 treatment. Having observed a rescue of most of the phenotypes associated with the homozygous duox mutants, we wondered whether T4 treatment also diminished the size of the thyroid gland in the duox mutants. Anecdotally, we had noted that one of the homozygous duox sa9892+/− mutant animals in the treated group had a small external goitre before treatment, but the goitre resorbed after 2 weeks of treatment. In comparison, a homozygous mutant sibling in the untreated group, that also had an external goitre, showed an increase in the size of the goitre during the course of the experiment (data not shown). This suggested that T4 treatment might lead to a diminution in the size of the thyroid glands in the homozygous mutant animals. To confirm whether this was the case, we sectioned and performed ISH for thyroglobulin on some of the treated and untreated homozygous mutant animals. We found that treatment led...
to a dramatic decrease in the thyroid hyperplasia normally associated with the duox homozygous mutants. However, some of the treated animals did retain some small areas of ectopic thyroglobulin staining in the head not seen in WT animals, suggesting that these animals had extensive ectopic thyroid follicular tissue prior to treatment (Fig. 7H–J).

**Methimazole phenocopies duox mutant phenotypes**

For final confirmation that the phenotypes found in the homozygous duox mutants were due to hypothyroidism, we asked whether exposure of WT fish to the goitrogen methimazole (1 mM) phenocopied the homozygous mutant phenotypes. To counter the influence of already circulating THs, we exposed the adult fish over a 3-month period. Treated animals became darker, owing to an increase in the number of melanophores (Fig. 8A–C). In addition, adult fish treated with methimazole failed to breed, as observed among homozygous duox mutant animals. They also developed external goitres (three out of seven) (Fig. 8F) and, internally, their thyroid follicles spread dramatically in area (Fig. 8G,H). This was reminiscent of the observations made in homozygous duox mutants (Fig. 8I). Finally, WT larval zebrafish continuously treated with methimazole from between 3 hpf and 5 hpf onwards showed resistance to follicular T4 immunostaining, similar to that found in duox sa9892−/− and duox sa13017−/− mutant larvae (Fig. 8D,E).
Fig. 3. Adult *duox* mutant zebrafish display an array of visible phenotypes. (A–C) 5× magnification of flank region showing the distribution of melanophores in WT, *sa9892*+/− and *sa989*−/− siblings. The apparent abundance of melanophores was statistically significant in *duox* mutants (D). Asterisks denote statistically significant differences (Bonferroni’s multiple comparisons test, ****P<0.0001). *duox* mutants also showed irregularities in stripe pattern in contrast to heterozygous siblings, shown here in a 2× magnification of the flank in *sa9892* siblings (E,F). Craniofacial anomalies were evident among mutants, with frontal height significantly shorter among mutants (G–I) (Bonferroni’s multiple comparisons test, *P<0.5, **P<0.01). Erythema in the thoracic region was prominent among mutants. This was especially noticeable in nacre backgrounds (L,N,O). *duox* mutants also suffered from perpetual fin damage, which manifest as ragged margins and tears (S–U). Scale bars: 1 mm.
DISCUSSION
The zebrafish has recently emerged as a new, genetically tractable model for investigating the molecular mechanisms underpinning thyroid organogenesis and function (Alt et al., 2006; Elsalini and Rohr, 2003; Guillot et al., 2016; McMenamin et al., 2014; Trubiroha et al., 2018; Wendl et al., 2002). Although a recent report described the larval phenotype associated with CRISPR generated bi-allelic loss-of-function duox mutations in F0 zebrafish (Trubiroha et al., 2018), there have been no prior reports describing the phenotypic consequences of fully characterised duox alleles in adult zebrafish. This is despite the fact that mutations in DUOX2 and DUOX1 have been shown to be associated with congenital hypothyroidism in humans for more than a decade (Aycan et al., 2017; Donkó et al., 2014; Jin et al., 2014; Johnson et al., 2007; Kizys et al., 2017; Tonacchera et al., 2009; Vigone et al., 2005).

Here, we describe a comprehensive assessment of the adult phenotypes associated with homozygosity of two loss-of-function alleles of zebrafish duox in adult fish. The additional round of genome duplication in teleost fish (Taylor et al., 2003) notwithstanding, there only exists a single orthologue of duox in zebrafish, instead of the two orthologues present in tetrapods (DUOX1 and DUOX2) (Kawahara et al., 2007). Remarkably then, in this instance, zebrafish has less genetic redundancy for this gene than is commonly found in this system. Thus, assessing phenotypes associated with homozygosity of the single duox orthologue in zebrafish has allowed us to model the effect of losing the function of both duox orthologues in tetrapods. This is particularly important as mutations in both DUOX1 and DUOX2 in humans have been associated with a more severe form of CH (Aycan et al., 2017), suggesting that DUOX1, while normally playing a minor role in TH synthesis in humans, does partially compensate for the loss of DUOX2 in humans.

Amongst the various adult phenotypes displayed by the homozygous duox mutants, most have been previously observed following pharmacological disruptions in thyroid hormone synthesis or in mutant strains where the hypothalamic–pituitary–thyroid (HPT) axis in zebrafish is affected. For example, goitrogen treatments, thyroid ablation and tshr mutant strains display alterations in pigmentation (McMenamin et al., 2014), similar to those we found in homozygous duox mutants. More specifically, thyroid ablated zebrafish have a darker striped pattern due to an increase in the density of melanophores within each stripe (McMenamin et al., 2014), akin to our homozygous duox mutants. Another notable phenotype in our homozygous mutants reminiscent of prior findings in goitrogen-treated zebrafish was erythema in the proximity of the operculum. Schmidt and Braunbeck (2011) came across a striking histopathological phenotype following treatment of WT zebrafish with the goitrogen, phenylthiouracil (PTU) (Elsalini and Rohr, 2003), wherein treatment resulted in excessive proliferation of blood vessels surrounding the thyroid follicles. This proliferation is attributed to hyperemia resulting from blood aggregation in proximally swollen blood vessels surrounding the thyroid follicles and is concentration-dependent, with the highest concentrations leading to hyperemia (Schmidt and Braunbeck, 2011). Macroscopically, this proliferation of vasculature manifests as erythema, giving a red colour to the entire opercular region. While we did not perform a histological examination of the vasculature, our macroscopic observations are consistent with these reported findings, with all our mutants displaying this conspicuous redness of the opercular region. The colouration was most notable amongst younger animals and especially apparent in backgrounds lacking melanophores. Space constraints together with follicular expansion and vascular proliferation in the pharyngeal region could also explain for the flaring opercula observed in some mutants, although this could also be due to thyroid hyperplasia, which was also noted by Schmidt and Braunbeck (2011) in their PTU-treated fish.

We were also able to induce this chronic hypothyroid/goitrogenic state in WT animals following treatment with methimazole, resulting in similar phenotypic outcomes. In our reverse experiment, however,
it was very interesting to note that while T₄ treatment of mutants resolved the goiters, some follicular staining remained in ectopic regions. Even so, the overall amount of thyroid tissue was largely diminished. It has been reported that at concentrations ≥25 mg/l of PTU, follicular encroachment is found in the gills of zebrafish, suggesting ectopic follicular expansion (Schmidt and Braunbeck, 2011). Follicular expansion following exposure to methimazole is attributed to thyroid hyperplasia, both, in zebrafish (Schmidt and Braunbeck, 2011) and in frog tadpoles (Hsu et al., 1974). This is regarded to be the first step in compensating for TH production via TSH (Schmidt and Braunbeck, 2011). Concentration dependent increases in the extent of follicular hypertrophy and hyperplasia have also been reported in the fathead minnow (*Pimephales promelas*), when exposed to the thyroid peroxidase inhibitor 2-mercaptobenzothiazole (Nelson et al., 2016). *duox* mutants and methimazole-treated WTs presented with amplified thyroglobulin...
expression that showed follicular crowding in the pharyngeal regions and invasion in other ectopic locations. Thyroid dyshormonogeneis (TD) is among the leading causes of CH, and ectopia (ectopic thyroid) is the commonest subtype of TD (De Felice and Di Lauro, 2004). Ectopic thyroid glands have recently been reported in human \( DUOX2 \) mutations wherein scintigraphy revealed submandibular and sublingual thyroid ectopic locations (Kizys et al., 2017).

Teleost fins have garnered interest within the scientific community not only due to their extensive morphological diversity, but also due to their remarkable regenerative capacities (Johnson and Bennett, 1999; Nakatani et al., 2007). Fins are composed of multiple branched and multi-segmented rays covered in a thin layer of epidermal cells. Individual rays consist of a pair of hemirays. Mature hemirays, known as lepidotrichia, are surrounded by a monolayer of osteoblasts that synthesise the bone matrix. With no musculature present, the remainder is made up of mesenchymal cells with nerve fibres and vasculature running along and inside the fin rays. Because of the constant growth, renewal and maintenance of the fins, it is relatively uncommon to find animals in aquaria with damaged fins (Wills et al., 2008). Thus, the ragged fins in the \( duox \) mutants stand out. We have found that the presence of ragged fins is ameliorated, however, by treatment of the mutants with \( T_4 \).

CH has been associated with cephalic and facial defects and developmental neurological abnormalities (Gamborino et al., 2001). Such defects have been attributed to improper development of the cranial neural crest (CNC), which is a transient population of migratory embryonic stem cells. Arising from the neural ectoderm, these cells contribute to a long list of cell types, including bone, cartilage, craniofacial connective tissue, corneal stroma and endothelium, iris stroma, ciliary body stroma and muscles, sclera and the trabecular meshwork of the eye (Barembaum and Bronner-Fraser, 2005; Minoux and Rijli, 2010). An investigation of craniofacial morphogenesis using rats exposed to methimazole revealed a 25% reduction in the overall head size throughout gestation (Gamborino et al., 2001). These findings are consistent with observations on craniofacial shape in zebrafish \( manet^{ppr23k} \) mutants as well as metronidazole-mediated thyroid ablated transgenics \( Tg(tg:nVenus-2a-nfnB)^{pp-r} \), which have narrower heads than controls (McMenamin et al., 2014). Further evidence on the role of THs has been gathered using pharmacological and morpholino-based approaches in zebrafish larvae. In one study, methimazole treatment resulted in reduced head depth and shorter jaw length (Liu and Chan, 2002). In another study, methimazole and PTU were found to inhibit pharyngeal arches and ceratohyal cartilage development, while knockdown of \( thraa \) (thyroid receptor \( \alpha \) a) led to malformations in the Meckel’s
and ceratohyal cartilages (Bohnsack and Kahana, 2013). Our observations of the shorter frontal height among duox sa9892−/−, sa13017−/− and sa9892/sa13017 animals are yet another indicator of TH deficiency.

Furthermore, duox mutants appear to experience several phenotypes associated with retarded growth and development. These include delayed growth rate, and delayed or incomplete swim bladder morphogenesis and barbel emergence. As development is

Fig. 7. T4 treatment alleviates phenotypic anomalies in duox mutants. T4-treated mutants show an improvement in fin health, compared to untreated mutants (A,B). Pigment changes are evident among T4-treated mutants. C−F show a 5× magnification of the distribution of melanophores on the flank region of sa9892+/− and sa9892−/− siblings, with a significant reduction in melanophore number (G). Asterisks denote statistically significant differences (Bonferroni’s multiple comparisons test, ****P<0.0001). Goitres resolve following T4 administration, but small ectopic thyroids are still evident (black arrowheads) (I,J). Scale bars: 1 mm.
underway, fish standard length (SL) is subject to both genetic and environmental factors, thus introducing variation amongst siblings. Indeed, environmental influences on SL is clearly apparent as larvae reared in groups show greater variation in SL than larvae raised individually (Parichy et al., 2009). SL is, thus, regarded as a more reliable measure of fish maturation than age (Parichy et al., 2009; Singleman and Holtzman, 2014). Considering the mean values for SL for our groups, it was clearly apparent that at 3 months of age all mutant groups were significantly shorter (i.e. less mature) than their heterozygous or WT siblings. Remarkably, by 6 months, however, the homozygous duox mutant fish caught up with the WT and heterozygotes siblings. This suggests that it is not growth per se, but the state of maturation, which is dependent on thyroid hormones. This finding is consistent with findings in non-metamorphosing Xenopus laevis tadpoles, which become giants and can live for years in an immature neotenic state (Rot-Nikcevic and Wassersug, 2004). This arrested development, associated with continued growth, has been attributed to a lack of thyroid glands in these animals (Rot-Nikcevic and Wassersug, 2004). In fish, definitive evidence of TH insufficiency causing metamorphic stasis is well appreciated from studies on flatfish. Larvae of the summer flounder (Paralichthys dentatus), when treated with thiourea, do not develop beyond early metamorphic climax (Schreiber and Specker, 1998). Likewise, olive flounder (Paralichthys olivaceus) larvae treated with the goitrogen, thiourea, enter metamorphic stasis and become giant larvae (Inui and Miwa, 1985). Although metamorphosis among the roundfish is less dramatic, several examples illustrate the dependence of metamorphosis on THs. Thiourea treatment was found to arrest metamorphosis in the coral trout grouper (Plectropomus leopardus) (Trijuno et al., 2002), orange-spotted grouper (Epinephelus coioides) (de Jesus et al., 1998) and the red sea bream (Pagrus major) (Hirata et al., 1989). Meanwhile, the pesticide chlorpyrifos, reported to cause reductions in serum concentrations of T4 and T3 (Slotkin et al., 2013), was recently found to prevent metamorphic completion in the convict surgeonfish (Acanthurus triostegus) (Holzer et al., 2017). In zebrafish, a 1 mM concentration of methimazole inhibited the larval to juvenile transition (Brown, 1997). However, larvae treated with a concentration of 0.3 mM eventually escaped the inhibition and continued development. While our duox mutants eventually reach normal adult size, this...
might be associated with an incomplete metamorphic or immature state. Alternatively, there may be some genetic redundancy present in zebrafish, whereby a different source of H$_2$O$_2$ in the thyroid follicles might be capable of partially compensating for the loss of duox function. Indeed, another NOX isoform, NOX4, has been described in human thyrocytes. Unlike Duox though, NOX4 generates H$_2$O$_2$ in the intracellular compartment (Weyemi et al., 2010). It may thus be important to generate double mutants for duox and nox4 to determine the contribution of Nox4 in thyroid hormogenesis.

In the zebrafish, swim bladder inflation is dependent on THs (Godfrey et al., 2017; Liu and Chan, 2002). The posterior chamber of the swim bladder inflates around 4.5 dpf while the anterior chamber inflates by 21 dpf (Winata et al., 2009). These events appear to coincide with peaks in whole body T$_3$ at 5 dpf and 10 dpf and T$_4$ at 21 dpf (Chang et al., 2012). Previously, it was found that swim bladder inflation was significantly delayed in thyroid-ablated zebrafish, where the anterior chamber of the bladder inflated ~50 dfp, compared to ~20 days in controls (McMenamin et al., 2014). Similar observations were also made in thyroid-ablated Danio albolineatus (McMenamin et al., 2014). There also exists sufficient evidence of how pharmacologically disrupted thyroid processes affect swim bladder inflation. Ecological assessments of aquatic pollutants often employ key morphological events during fish development as predictive approaches. 2-Mercaptobenzothiazole (MBT), commonly used for rubber vulcanization, is found to occur in environmental water bodies. MBT is a potent TPO inhibitor and its role was recently examined in swim bladder inflation in the fathead minnow (Pimephales promelas) (Nelson et al., 2016) and zebrafish (Stinckens et al., 2016). Among minnows, larvae continuously exposed to MBT showed a concentration-dependent decrease in anterior lobe size (Nelson et al., 2016). Meanwhile, MBT-treated zebrafish larvae were reported to fare worse than minnows, where 22% of larvae exposed to the highest concentration failed to inflate the anterior chamber (Stinckens et al., 2016). Interestingly, even though both species belong to the order Cypriniformes, a compensatory T$_4$ response has been reported in the fathead minnow at 21 dpf (Nelson et al., 2016) but not in the zebrafish (Stinckens et al., 2016), suggesting species-specific differences. Our homozygous duox mutant animals also displayed a significant delay in anterior chamber inflation, suggesting that Duox is essential for this process, likely through its role in thyroid hormone synthesis.

Barbels are yet another easily observable phenotypic trait influenced by thyroid hormones. In zebrafish, both pairs develop as epithelial buds around 30–40 dpf, following the emergence of pelvic fin rays (Hansen et al., 2002; Parichy et al., 2009). Thus far, only one study has reported barbel emergence to be influenced by THs. Thyroid ablation via Mzt of Tg(tg:nVenus-2a-nfnB) of Danio rerio and D. albolineatus resulted in the absence of sensory barbels (McMenamin et al., 2014). Similarly, manet mutants were also found to lack barbels (McMenamin et al., 2014). Our homozygous duox mutants also show impairment of barbel emergence, consistent with their hypothyroid state. However, it is notable that a subset of the sa8992^-/- mutations eventually did grow barbels, similar to their body length catch-up phenotype.

In humans, thyroid dysfunction during pregnancy has been positively associated with adverse maternal/foetal outcomes, including infertility, miscarriage, pre-eclampsia, pre-term (before 37 weeks) birth and maternal thyroid dysfunction postpartum (Hernández et al., 2018; Stagnaro-Green et al., 2011; Velasco and Taylor, 2018). TH is essential for early development and maturation of the foetal brain and maternal transfer of TH is especially important during the first trimester (Cooper and Biondi, 2012) since the embryo does not begin synthesising THs until 12–13 weeks into gestation (Casey and Leveno, 2006). The British Thyroid Foundation suggests prescribing levothyroxine to hypothyroid women trying to conceive in order to address these negative consequences of hypothyroidism on fertility and pregnancy. Intriguingly, we also noted significant defects in fertility in both sexes in our homozygous duox mutants. Although we do not currently know the reason for infertility in the duox mutants, a potential cause may be due to failure in mating behaviour as a consequence of the observed effects on pigmentation in the mutants. It has previously been noted that, in zebrafish, both sexes experience diurnal changes in their stripes and interstripe colours, a process termed ephemeral sexual dichromatism, during mating and spawning (Hutter et al., 2012). Another study reported that females utilise yellow colouration for sex recognition (Hutter et al., 2011). This ties in well with xanthophore deficiency reported in thyroid ablated, hypothyroid zebrafish (McMenamin et al., 2014), and by extension, the duox mutants. However, it is notable that casper strains of zebrafish, which lack xanthophores altogether, can successfully breed (White et al., 2008). Thus, there may be additional factors that may be contributing to infertility in duox mutants.

Associations between thyroid status and reproduction in teleosts have been previously reviewed (Cyr and Eales, 1996). Four physiological pre-requisites have been recognised as essential to spawning behaviour and fertility in fish: (1) the completion of vitellogenesis in the ovaries, (2) maturation and ovulation of oocytes stimulated by pituitary luteinizing hormone (LH), (3) completion of spermatogenesis, and (4) sufficient production and storage of milt (semenal plasma and mature sperm) in the sperm duct. These are largely regulated by the endocrine system (Kobayashi et al., 2002). T$_3$ enhances the response of the ovarian follicles to gonadotropins, thus facilitating secretion of 17β estradiol (Cyr and Eales, 1988). This regulates the production of vitellogenin by the liver, and in studies on Great Lakes salmonids it has been suggested that lowered T$_3$ levels may impair oocyte production (Leatherland and Barrett, 1993). In the fathead minnow (Pimephales promelas), Methimazole treatment led to a reduction of the cortical alveolus oocytes, relative to control females. Meanwhile, in post-spawning males, control animals showed an increase in the number of spermatozoa and a decrease in the number of spermatogonia. This increase in spermatoza was not observed in methimazole-treated cohorts, suggesting that hypothyroidism affects spermatogenesis (Lema et al., 2009). Among the African sharp-tooth catfish (Clarias gariepinus), pre-spawning males treated with thiourea were shown to have narrower seminiferous tubules and fewer spermatozoa was not observed in methimazole-treated cohorts, suggesting that hypothyroidism affects spermatogenesis (Lema et al., 2009). Intriguingly, hypothyroidism in humans has also been associated with impaired spermatogenesis and sperm abnormalities (La Vignera and Vita, 2018). We have found that fertility in our homozygous duox mutants can be restored in both sexes and we can successfully raise offspring to adulthood from a cross between a mutant male and WT female. This is in line with previous observations on growth-retarded (grt) mice. grt mice have autosomal recessive hypothyroidism, with females suffering lifelong infertility and males gradually acquiring fertility. When treated with THs, grt females showed an increase in the size of their uteri and ovaries, which was comparable with heterozygous and WTs. Furthermore, they engaged in copulatory behaviour and were able to conceive and deliver pups (Hosoda et al., 2008). Zebrafish duox mutants thus provide an excellent model to investigate the consequences of human CH associated with mutations in DUOX1 and DUOX2, and the mechanisms by which treatment with THs, even in adults, can
restore many of the defects caused by chronic hypothyroidism, including restoration of fertility in both males and females.

**Conclusion**

Overall, we found that homozygous mutants display a number of phenotypes, which can be ascribed to hypothyroidism, including growth retardation, pigmentation defects, ragged fins, thyroid hyperplasia and external goitre. By and large, the growth retardation defect is not permanent, as fish continue to grow despite being chronically hypothyroid, and ultimately catch up with their euthyroid heterozygous and WT siblings. This contrasts with findings in humans suffering from hypothyroidism, who remain growth retarded unless T4 treatment is initiated within weeks after birth. Most other phenotypes associated with chronic hypothyroidism in the duox mutant fish were rescued by T4 treatment, even if supplementation was not initiated until adulthood. These include recovery of fertility, return to normal pigmentation, improvement in fin morphology and return to normal size thyroid glands. In summary, duox mutant zebrafish provide a new and potentially powerful system to understand the consequences of chronic congenital hypothyroidism on growth and maintenance of body physiology, as well as the mechanisms of recovery of normal physiology following thyroid hormone supplementation. Thus, our duox mutant fish appear to be in a chronic hypothyroid/goitrogenic state, as indicated by their external goitres as well as internal expansion of thyroglobulin expressing tissue.

**MATERIAL AND METHODS**

**Ethics statement**

All experiments involving animals were approved by the local ethics committee and the Home Office.

**Animals and husbandry**

Adults and larvae were used in this study. The zebrafish (*D. rerio*) WT line used was AB. Mutant lines used were *duox sa9892* and *duox sa13017* (Kettleborough et al., 2013) and were obtained from the European Zebrafish Resource Center (EZRC). Compound heterozygotes for these mutant alleles were generated in-house. Both duox alleles were also crossed into nacre (*nac*) (Lister et al., 1999) and casper (White et al., 2008) strains for visualising larval thyroid follicles, swim bladder and adult erythema. In all cases, embryos were raised in sea salts (Sigma-Aldrich, 59883) medium containing 0.0001% Methylene Blue until 5 days post-fertilisation (dpf) and then transferred to the system where they were maintained at a temperature of 28°C, pH 7.4, constant salinity and a 14:10 photoperiod.

**PCR and genotyping**

Genomic DNA was extracted from caudal fin clips or whole larvae using lysis buffer, in a thermal cycler. The conditions for this procedure were 2 h at 55°C, 10 min at 95°C and a hold (if necessary) at 12°C. PCR was performed using ExTaq DNA polymerase (Takara RR001A) with the following primer pairs: for the *duox sa9892* allele, forward 5′-AGCAGGTACAACTCAAGCGT-3′ and reverse 5′-GACGTTCAAAGCAGAAGTGC-3′; for the *duox sa13017* allele, forward 5′-TGTTACACATTGAGGATGGA-3′ and reverse 5′-ACACCCACATAGGCTC-3′. PCR conditions were as follows: 36 cycles at 94°C for 30 s, 55°C for 30 s and 72°C for 30 s. Samples were subject to Sanger sequencing (GATC Biotech). Sequencing primers used were 5′-CTTGTGTCGCTTTGAGGAAATG-3′ for the *duox sa9892* allele and 5′-GTGACTCAAGTACAGGCGTCC-3′ for the *duox sa13017* allele. Siblings were stage-matched, phenotypically WT, heterozygous and homozygous animals obtained by crossing heterozygous carriers.

**Whole mount immunofluorescence**

Zebrafish larvae, at 5 dpf, were fixed overnight in 4% phosphate-buffered paraformaldehyde (PFA) (Sigma-Aldrich), at 4°C. This was followed by 15 min of dehydration in 100% methanol. Larvae were then transferred to fresh 100% methanol and stored at −20°C until usage. Larvae were gradually rehydrated to PBT, treated with 10 µg/ml proteinase K (Roche) for 30 min, briefly rinsed in PBT, and postfixed in 4% PFA for 20 min. Following further rinsing in PBT, larvae were immersed in blocking buffer (PBT containing 1% dimethylsulfoxide, 1% BSA, 5% horse serum and 0.8% Triton X-100) for 2 h. This was followed by overnight incubation, at 4°C, in blocking buffer containing the primary antibody (1:1000) against thyroxine (T4) (Bioryt orb11479). Overnight incubation was followed by several wash steps in PBT containing 1% BSA. Larvae were then incubated overnight, at 4°C, in blocking buffer containing the secondary antibody (1:250) Alexa Fluor 568 (Invitrogen A11057) (Optiz et al., 2011). Stained larvae were washed in PBT, imaged and then subject to genotyping.

**Histology and ISH**

*In situ* hybridization on sections of adult zebrafish was performed essentially as described (Paul et al., 2016). Briefly, adult zebrafish were fixed whole in 4% PFA for 1 week followed by a decalcification step in 20% EDTA, for 10 days. Animals were then cut at the operculum and mid trunk level and processed in a Leica TP1050 tissue processor in preparation for paraffin embedding. The embedding station used was a Leica EG1150H. The cut face of the tissue was oriented towards the leading edge of the paraffin block and sectioned at 5 µm thickness on a LeicaRM2255 microtome. Sections were arranged and held on Superfrost Plus™ slides (Thermo Fisher Scientific). Alternating sections were then taken forward for Haematoxylin and Eosin (H&E) staining and ISH. Sections were put through H&E staining via a Varistain 24-4 carousel (Thermo Fisher Scientific, Shandon). *Thyroglobulin* (tg) cDNA used for riboprobe synthesis was amplified using forward 5′-AGGTGGAGAAAATGTGTGGT-3′ and reverse 5′-CTCACAACGTGCGC-AATGACT-3′ primers. Digoxigenin-labelled probes were synthesised *in vitro* using a MEGAscript™ kit (Ambion).

**Body length and melanophore counts**

For measuring body length, all fish were briefly anaesthetised in 0.02% MS 222 (tricaine) (Sigma-Aldrich). They were then transferred onto an agarose bed in a petri dish and imaged at 0.73× magnification. For each fish two or three images were captured in order to include the entire length including the caudal fin. These part images were stitched together in Adobe® Photoshop to obtain a single image. The ‘ruler’ tool and ‘analyse measurement’ command in Photoshop CS5 were used on these images to calculate the length from the tip of the mandible to the caudal peduncle. For determining melanophore density, all fish were treated with epinephrine to contract pigment granules. To obtain a 1 mg/ml working solution, 0.1 g of epinephrine was dissolved into 100 ml of a 0.01% tricaine solution. Epinephrine is only partially soluble in water and thus, the solution was filtered to obtain a clear filtrate. The solution turns pink-orange during filtration. Fish were treated for 5 min in this working solution. They were then transferred onto an agarose bed in a petri dish and imaged at two locations on the lowest continuous stripe extending from the operculum to the trunk. A 5× magnification was used. The ‘multi-point’ tool on Fiji was used to manually count all melanophores contained within a stripe, in the field of view, for each image. All images were acquired on a Leica MZ16FA fluorescence stereomicroscope with a DC490 camera.

**Pharmacological treatments**

For rescued mutant phenotypes, a 12-week treatment with T4 (Sigma-Aldrich) was sustained in a closed system that closely resembled aquarium conditions. Four groups—sa9892+/− untreated, sa9892+/− treated, sa9892+/− untreated and sa9892+/− treated were subject to this regime, with each group comprising of four adult fish. T4 was added tri-weekly, at a concentration of 30 nM. Water was changed three times each week.

For phenocopy experiments, a 12-week treatment with methimazole (Sigma-Aldrich) was administrated, once again, in a closed system simulating aquarium conditions. This regime was applied to six WT adult fish, while six untreated WT animals comprised the control group. Methimazole was added tri-weekly, at a concentration of 1 mM. Water was changed three times each week.
For immunostaining, WT larvae were treated with methanazol (0.3 mM) from 0 dpf. Animals were then fixed at 5 dpf and stained for T\textsubscript{4} as described above.

**Statistical analyses**

GraphPad Prism 7 was used for statistical testing. Column statistics and analyses of variance were implemented for all data sets. For column statistics, we calculated median, s.d., s.e.m., confidence intervals and Gaussian distribution. The D’Agostino and Pearson test was used to check for Gaussian distribution. One-way ordinary ANOVA was used to analyse variance. Differences were considered significant at $P<0.0001$. Bonferroni’s multiple comparisons test was used to compare means between groups.

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**Competing interests**

The authors declare no competing or financial interests.

**Author contributions**

Conceptualisation: K.C., S.I., E.A.; Methodology: K.C., S.I.; Validation: K.C.; Formal analysis of origin and growth of the thyroid gland in zebrafish. We would like to thank Peter Walker, Grace Bako and Natalie Partington, at the Core Histology Facility, for their help with histological sectioning and staining. We would also like to thank the aquarium staff in the BSF unit for their care and support of the fish. We also extend our gratitude to Simone Costagliola and Pierre Gillotay, Université Libre de Bruxelles, Belgium, for their advice with the T\textsubscript{4} antibody protocols in larvae.

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