Gestational stress induces the unfolded protein response, resulting in heart defects

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ABSTRACT

Congenital heart disease (CHD) is an enigma. It is the most common human birth defect and yet, even with the application of modern genetic and genomic technologies, only a minority of cases can be explained genetically. This is because environmental stressors also cause CHD. Here we propose a plausible non-genetic mechanism for induction of CHD by environmental stressors. We show that exposure of mouse embryos to short-term gestational hypoxia induces the most common types of heart defect. This is mediated by the rapid induction of the unfolded protein response (UPR), which profoundly reduces FGF signaling in cardiac progenitor cells of the second heart field. Thus, UPR activation during human pregnancy might be a common cause of CHD. Our findings have far-reaching consequences because the UPR is activated by a myriad of environmental or pathophysiological conditions. Ultimately, our discovery could lead to preventative strategies to reduce the incidence of human CHD.

KEY WORDS: FGF signaling, Heart development, Hypoxia, Unfolded protein response

INTRODUCTION

Congenital heart disease (CHD) is an abnormality in the structure or function of the heart that arises before birth. It is the most common form of human birth defect, with an incidence of ~9 per 1000 live births (van der Linde et al., 2011). The etiology of CHD is complex, with both genetic and environmental factors playing a role. Genetic factors have been widely investigated (reviewed by Andersen et al., 2014; Fahed et al., 2013). However, only ~20% of CHD cases can be attributed to a specific genetic cause (Blue et al., 2012). Thus, non-genetic factors are likely to be important in human heart development. This idea is supported by the large phenotypic variability seen between individuals with the same genetic mutation (Schott et al., 1998; Benson et al., 1999), and this is also true with variable heart defects in inbred mice (Abu-Issa et al., 2002). Epidemiological studies have identified significant environmental risk factors for human CHD, including maternal exposures to alcohol, isoretinoin, thalidomide, antiseizure medications, antiretroviral medications, environmental teratogens and infectious agents. Embryo hypoxia is another well-recognized risk factor caused by smoking, living at high altitude, maternal diabetes, high body mass index, hypertension or prescription medications (Watkins et al., 2003; Jenkins et al., 2007; Zheng et al., 2013; Webster et al., 2014; Ornoy et al., 2015; Ramakrishnan et al., 2015; Sullivan et al., 2015). The specific effects of environment on embryonic development have been studied by exposing pregnant animals to a variety of conditions. Hypoxia has been studied in this way for almost 200 years, causing a variety of birth defects, including heart defects (Geoffroy Saint Hilaire, 1820; Ingalls et al., 1952; Sparrow et al., 2012). We have recently shown that short-term gestational hypoxia significantly increases the penetrance and severity of vertebral defects in mouse models of congenital scoliosis (Sparrow et al., 2012). At the molecular level, hypoxia disrupts embryonic FGF signaling, leading to a temporary failure of somitogenesis. Heart formation in vertebrates has been well studied (reviewed by Miguerol and Kelly, 2013). The linear heart tube forms from the first heart field (FHF). These cells contribute mainly to the left ventricle and both atria of the mature heart. The linear heart tube elongates and loops to the right, driven by the addition of second heart field (SHF) cells to the arterial and venous poles. These cells contribute mostly to the right ventricle and inflow/outflow regions of the heart. The proliferation of SHF cells is controlled by FGF and β-catenin/WNT signaling, and their subsequent differentiation is controlled by BMP and non-canonical WNT pathways. Perturbation of proliferation, or premature activation of differentiation or apoptosis, for example by conditional deletion of FGF signaling components in the SHF, results in OFT elongation, alignment and septation defects (Hutson et al., 2006; Ilaqan et al., 2006; Park et al., 2006). In humans such defects include membranous ventricular septal defects (VSDs), persistent truncus arteriosus (PTA), double outlet right ventricle (DORV), transposition of the great arteries (TGA), overriding aorta (OA) and tetralogy of Fallot (TOF), and account for as much as 30-60% of all human CHD (Thom et al., 2006; Bruneau, 2008).

Here we present evidence that short-term gestational hypoxia can perturb mammalian heart formation, resulting in defects of OFT alignment and/or elongation. We show that hypoxic exposure results in reduced cellular proliferation in the SHF and elsewhere. In the SHF this is likely to be due to a downregulation of FGF signaling, mediated by the specific loss of the receptor FGFR1. We show that hypoxia induces the unfolded protein response, causing a global downregulation of protein translation. Since FGFR1 protein has a short half-life (Haugsten et al., 2005), this results in rapidly reduced receptor and signaling levels. Our findings might provide a mechanism for the genesis of some of the 80% of cases of human CHD.
CHD that remain unexplained by a purely genetic etiology. This mechanism is also likely to apply to birth defects in addition to CHD, as FGF signaling is required for numerous processes during embryogenesis.

RESULTS

In utero exposure of mouse embryos to lowered oxygen levels perturbs cardiac outflow tract formation

We investigated if exposure of mouse embryos to hypoxia in utero induces heart defects. Previous studies exposing mice to hypoxia (Ingalls et al., 1952) induced a low rate of VSDs. We performed similar experiments using more extreme hypoxia to maximize the rate of induced heart defects. Mice can tolerate as low as 5.5% oxygen at normal atmospheric pressure (Sparrow et al., 2012). We therefore exposed pregnant C57BL/6J mice to 5.5% oxygen for 8 h (hypoxia), and then returned them to room air (normoxia) until embryonic day (E) 17.5, when embryonic heart morphology was examined (Table 1). Groups of pregnant mice were exposed on different days encompassing heart development (E7.5-10.5). Heart defects were observed in some embryos exposed from E8.5-10.5, with a statistically significant peak at E9.5. At this stage, 21/48 embryos had heart defects. By contrast, low oxygen exposure did not significantly increase the incidence of embryo death, except at E9.5. All abnormal embryos exposed at E9.5 (21/21) had defects typical of a disruption of outflow tract (OFT) alignment or elongation (VSD, OA, DORV, TGA), while some had additional defects (Table S1). Thus, low oxygen exposure perturbed cardiac OFT alignment development with maximum effect at E9.5.

Alteration in cardiac OFT phenotype is apparent from E10.5

To determine the developmental origin of the cardiac defects, we examined OFT morphology in embryos exposed to 5.5% oxygen at E9.5 for 8 h and then allowed to develop in normoxia for a further 16-24 h. Control and exposed embryos were matched for somite number (range 33-38), and the OFT myocardium visualized by whole-mount myosin heavy chain (MHC) immunostaining. Optical projection tomography (OPT) was used to construct three-dimensional representations of the OFT myocardium (Fig. 1A,B). The majority of exposed embryos had altered OFT morphology, with a shorter distal OFT and an altered angle between proximal and distal OFT. There was a strong positive correlation between these parameters for control embryos (Fig. 1C, black points, \( P=0.0086 \)) but not for exposed embryos (Fig. 1C, red points, \( P=0.3221 \)). Thus, hypoxia-induced cardiac defects may result from a perturbation of distal OFT formation.

Low oxygen exposure affects proliferation in the SHF

The OFT forms primarily from cells of two embryonic lineages: cardiac progenitors of the SHF and cardiac neural crest cells (reviewed by Kelly, 2012; Neeb et al., 2013). Disruption of SHF development can result in defective OFT elongation, alignment and/or septation. Such defects can arise from activated apoptosis (Ilagan et al., 2006), premature differentiation (Liao et al., 2008) or reduced cell proliferation (Xu et al., 2004; Ilagan et al., 2006). No apoptosis was detected by active/pro-caspase 3 or TUNEL staining in the SHF, OFT or myocardium of control or exposed embryos (Fig. 1D,E, Fig. S1A-T). However, increased numbers of apoptotic cells were present in the neuroepithelium of the posterior part of the neural tube, otic vesicle and gut of exposed embryos, confirming that we were able to detect apoptotic cells by these techniques (Fig. S1A-T). Likewise, we did not detect any change in expression levels or location of the myocyte differentiation marker MHC between control and exposed embryos (Fig. 1G,H). By contrast, there was a clear reduction in the extent of proliferation in the SHF, OFT myocardium, branchial arch and forebrain in exposed embryos (Fig. 1J-L, Fig. S1U-Z,A-C). Quantification of the percentage of phosphorylated histone H3-positive nuclei indicated that proliferation in the SHF and forebrain was more affected than that in the myocardium or branchial arch. Thus, low oxygen exposure results in a significant reduction of SHF cell proliferation, but does not induce apoptosis or premature differentiation.

Low oxygen exposure reduces FGF signaling in the SHF

We have shown that low oxygen exposure in utero reduces FGF signaling in the presomitic mesoderm (PSM) (Sparrow et al., 2012). In the SHF, reduction or loss of FGF signaling causes reduced cell proliferation and OFT defects (Hutson et al., 2006; Park et al., 2006, 2008; Zhang et al., 2008). We therefore examined the expression of direct target genes of FGF signaling in the SHF. Spry1, Spry2 and Spry4 transcripts in the SHF were significantly reduced after 8 h exposure in the SHF (Fig. 2A-C, Fig. S2N-Q). To gauge the dynamics of FGF signaling disruption, we exposed embryos for 1, 2, 4 or 8 h to 5.5% oxygen and then examined Spry2 transcript levels. The significant reduction in transcript levels in the SHF was only observed in embryos exposed for 8 h (Fig. S2A-F).

We next examined a more upstream component of FGF signal transduction. Depending on cellular context, FGF signal transduction can occur through three pathways: MAPK (ERK), AKT and PKCγ (Dorey and Amaya, 2010). ERK is the main pathway used by cells of the SHF (Ilagan et al., 2006; Zhang et al., 2008). Phosphorylated ERK1/2 (also known as MAPK3/1) was significantly reduced in the SHF after 8 h exposure (Fig. 2D-F). To investigate the dynamics of phosphorylated ERK1/2 loss, levels were examined after 0.5, 1, 2, 4 or 8 h exposure (Fig. S2G-M). As little as 1 h exposure caused a significant loss of phosphorylated ERK1/2. Thus, the reduced proliferation in SHF cardiac progenitor cells is likely to be due to reduced FGF signaling, and this occurs extremely rapidly upon exposure to hypoxia.

Table 1. In utero low oxygen exposure induces heart defects

<table>
<thead>
<tr>
<th>Exposure stage</th>
<th>Heart morphology</th>
<th>Abnormal (%)</th>
<th>P-value</th>
<th>Resorbed (%)</th>
<th>P-value</th>
</tr>
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<tbody>
<tr>
<td>Control</td>
<td>Normal</td>
<td>31</td>
<td>6.1</td>
<td></td>
<td>8.3</td>
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<tr>
<td>E7.5</td>
<td>27</td>
<td>2</td>
<td>0</td>
<td>0.30</td>
<td>3.8</td>
</tr>
<tr>
<td>E8.5</td>
<td>33</td>
<td>7</td>
<td>17.5</td>
<td>0.13</td>
<td>8.9</td>
</tr>
<tr>
<td>E9.5</td>
<td>27</td>
<td>21</td>
<td>43.8</td>
<td>0.0001</td>
<td>11.9</td>
</tr>
<tr>
<td>E10.5</td>
<td>29</td>
<td>7</td>
<td>19.4</td>
<td>0.10</td>
<td>33.9</td>
</tr>
</tbody>
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\( P \)-values were obtained by one-tailed Fisher’s exact test.
reduced directly in response to cellular hypoxia. FGF signaling in
the SHF involves the ligands FGF8 (Brown et al., 2004; Ilagan et al.,
2006; Park et al., 2006) and FGF10 (Marguerie et al., 2006;
Watanabe et al., 2010) and the receptors FGFR1 and FGFR2
(Marguerie et al., 2006; Park et al., 2008; Zhang et al., 2008). We
focused on FGF8 and FGFR1, as these are prominent in SHF
formation (Marguerie et al., 2006; Park et al., 2008; Watanabe et al.,
2010). Levels of Fgf8 transcripts were significantly reduced in the

Fig. 1. Effects of low oxygen exposure of E9.5 mouse embryos on OFT morphology and on apoptosis, differentiation and proliferation of SHF cells.

(A-C) Comparison of three-dimensional reconstructions of control (n=11) and hypoxia-exposed (8 h, n=11) E10.5 OFT generated by optical projection
tomography (OPT). Representative images are shown of OFT from control (A) and exposed (B) embryos. Yellow lines indicate the measured distal OFT length,
and black lines indicate the angle measured between the distal and proximal OFT. (C) The relationship between distal OFT length and angle between distal and
proximal OFT at E10.5 for control (black dots) and exposed (red dots) embryos. The measurements for the images shown in A and B are indicated by open circles
in black and red, respectively. Spearman correlation P-values were 0.0086 (control) and 0.3221 (exposed). (D-L) Comparison of protein levels in control and
exposed E9.5 mouse embryos using immunohistochemistry on sagittal paraffin sections. (D,E) Comparison of active/pro-caspase 3 levels (n=10 control and
n=10 exposed, red), (G,H) myosin heavy chain (MHC, n=10 control and n=19 exposed, green) and (J,K) phospho-histone H3 (n=10 control and n=10 exposed, green).
Sections were counterstained with Hematoxylin (D,E, light blue) or TO-PRO-3 (G,H,J,K, red). For clarity, in J,K the brightness levels in the green channel only
were increased. (F,I) Diagrams indicating the relative positions of the SHF (dark blue), pharyngeal endoderm (green), OFT (light blue) and left ventricle (V, red)
and left atrium (A, orange) in a sagittal section of an E9.5 embryo. (L) Quantification of percentage phospho-histone H3-positive nuclei. Red dots indicate
quantification of the images shown in J and K. ***P<0.001. For sections, rostral is to the top and caudal to the bottom. Location of the SHF is indicated by brackets.
Scale bar in K: 80 μm in A,B,J,K; 130 μm in D,E; 140 μm in G,H. See also Fig. S1.
SHF from 4 h exposure, but not after shorter exposures (Fig. 3A-C, Fig. S3A-F). Thus, Fgf8 transcript levels were not reduced until well after phosphorylated ERK1/2 levels were significantly reduced (4 h compared with 0.5 h). This suggests that the Fgf8 gene might itself be a target of FGF signaling, and that its reduced transcription is a consequence, rather than the cause, of reduced FGF signaling. This is supported by the observation that conditional knockout of Fgfr1 in the midbrain-hindbrain boundary greatly reduces FGF signaling and abolishes Fgf8 expression (Jukkola et al., 2006).

We next examined FGFR1 protein expression. In control embryos, FGFR1 protein was concentrated in perinuclear cytoplasmic puncta, with lower levels of background staining throughout the cell (Fig. 3D). This is consistent with previous observations in different cell types in vivo and in vitro (Prudovsky et al., 1994; Ozaki et al., 2000). The puncta co-stained with the cis-Golgi protein GM130 (also known as GOLGA2) (Fig. 3E,F). By contrast, after 8 h exposure there was a reduction in both total FGFR1 (7/10 embryos, Fig. 3G) and Golgi-localized FGFR1 (8/10 embryos, Fig. 3K). Concomitantly, Golgi subcellular localization did not appear to be affected (Fig. 3E,F,I,J). Low oxygen exposure had a rapid effect on FGFR1 protein expression, with exposure for as little as 1 h resulting in a significant loss of Golgi-localized FGFR1 protein and total FGFR1 significantly reduced after 4 h exposure (Fig. S3G-T). We confirmed this reduction by immunoblot of whole-embryo extracts (Fig. 3O, Fig. S3B′). Thus, FGFR1 protein levels are rapidly reduced in the SHF in response to low oxygen exposure, and there is some specificity to this response.

**SHF cells in embryos exposed to lowered oxygen levels in utero are hypoxic**

We next investigated how hypoxia causes FGFR1 protein loss. Maternal low oxygen exposure is likely to have many effects on the embryo. Previously, we have shown that the embryonic myocardium is particularly susceptible to low oxygen exposure (O’Reilly et al., 2014). We therefore investigated whether the SHF was similarly affected. We measured embryonic cellular oxygen levels using the compound Hypoxyprobe (Raleigh et al., 1999). Pregnant mice were injected with Hypoxyprobe and then exposed to 5.5% oxygen for 3 h. Limited staining was detected in control embryos under these conditions (Fig. 4A,B). By contrast, there was strong staining throughout exposed embryos, including the SHF (Fig. 4C,D). The response to cellular hypoxia in mammalian embryos is mediated by hypoxia inducible factor (HIF1) (reviewed by Dunwoodie, 2009). The HIF1α subunit of this transcription factor is rapidly degraded in normoxia (Wang et al., 1995), but it is stabilized and transported to the nucleus when cellular oxygen levels fall below 2% (Jiang et al., 1996). Nuclear HIF1α staining was detected in the myocardium of embryos after as little as 0.5 h exposure (Fig. 4E,F,H, Fig. S4A-H,M) (O’Reilly et al., 2014). Nuclear staining was also visible in the gut (Fig. S4N). No nuclear staining of HIF1α was visible in the SHF of embryos exposed to low oxygen (Fig. 4E-G), despite these cells being hypoxic, although there was a slight increase in the total amount of HIF1α staining.
SHF cells in embryos exposed to lowered oxygen levels in utero induce the unfolded protein response

One cellular consequence of hypoxia is disruption of protein folding in the endoplasmic reticulum (ER) (Feldman et al., 2005). If such disruption is severe, the unfolded protein response (UPR) will be induced. This is a coordinated cellular program that reduces the rate of protein synthesis generally, increases the expression of molecular chaperones to assist protein refolding,
upregulates the ER-associated degradation (ERAD) pathway to degrade terminally misfolded proteins, and induces apoptosis if ER stress persists (Kitamura, 2013).

We examined the expression of a late marker of the UPR, DNA damage-inducible transcript 3 (DDIT3, also known as CHOP) to determine if the UPR is induced in low oxygen-exposed embryos. DDIT3 was robustly induced in the SHF of 8/12 embryos after 8 h exposure, and in a subset of exposed embryos from as early as 2 h (Fig. 4I-K, Fig. S4O-U). We also analyzed the production of an immediate early marker of the UPR, namely spliced $Xbp1$ transcript. Significant levels of spliced $Xbp1$ were detectable by qPCR in whole-embryo RNA extracts after as little as 0.5 h exposure (Fig. 4L). This suggests that the UPR is rapidly induced in embryos exposed to low oxygen levels. Thus, the specific loss of FGFR1 protein expression in embryos exposed to low oxygen levels correlates with both hypoxia and induction of the UPR in the SHF.

FGFR1 protein levels are reduced in vitro in response to hypoxia and other cellular stressors

We next developed an in vitro model system to investigate the molecular mechanism by which FGFR1 is lost in response to hypoxia. We used mouse muscle satellite C2C12 cells because they express FGFR1 and can activate the FGF pathway through ERK (Erck et al., 1998). Cells were cultured for 1-4 h in 0.1% oxygen and then protein extracts analyzed by immunoblot (Fig. 5A,B). In keeping with our in vivo results, levels of FGFR1 and phosphorylated ERK1/2 were significantly reduced by hypoxia, whereas total ERK1/2 and β-tubulin levels remained constant. The degree of reduction was similar to that observed in immunoblots of whole embryos exposed to 5.5% oxygen for 1-4 h (Fig. 3O). Exposed cells also had significantly increased levels of the translation initiation factor phosphorylated EIF2α and DDIT3, indicating robust UPR activation (Fig. 5A,B).
We also tested the effects of other UPR-inducing stressors on FGFR1 stability in vitro. Short-term treatment of cells with hydrogen peroxide, increased temperature or high pH all reduced FGFR1 protein levels by a similar extent to hypoxic treatment (Fig. S5A-C). Phosphorylated EIF2α was induced in all cases, confirming UPR activation (data not shown). Thus, other cellular stressors that induce the UPR, including increased reactive oxygen species or hyperthermia, also result in reduced FGFR1 protein levels in vitro.

**FGFR1 protein levels are reduced in vitro and in vivo due to UPR-triggered translation inhibition**

The UPR involves three separate signaling pathways mediated by IRE1 (also known as ERN1), PERK (also known as EIF2AK3) and ATF6. We next investigated whether the reduction in FGFR1 protein levels in vitro is due to the action of one particular pathway. The UPR activates the IRE1 pathway via Xbp1 transcript splicing, enabling translation of this transcription factor, increasing the expression of chaperones and activating the degradation of misfolded proteins. We compared FGFR1 protein levels in hypoxic cells with or without treatment with a small molecule inhibitor of IRE1, 4-methyl umbelliferone 8-carbaldehyde (4µ8c) (Axten et al., 2012). Hypoxia caused a significant reduction in FGFR1 protein levels, and treatment with 4µ8c did not alter this response (Fig. 6A). To confirm the efficacy of 4µ8c treatment, we monitored Xbp1 splicing levels using the transiently transfected FLAG-XBP1ΔDBD-venus reporter (Iwawaki et al., 2004). As expected, hypoxia activated Xbp1 splicing, and treatment with 4µ8c under hypoxic conditions prevented Xbp1 splicing (Fig. 6B). We also confirmed the specificity of 4µ8c action, as induced levels of phosphorylated EIF2α and DDIT3 under hypoxic conditions were unaffected by 4µ8c treatment (Fig. 6C,D).

The UPR also causes a global reduction in cap-dependent translation. This process is controlled by activation of the serine-threonine kinase PERK, which phosphorylates and inactivates the translation initiation factor EIF2α (Kitamura, 2013). We compared FGFR1 and phosphorylated ERK1/2 protein levels in hypoxic cells with or without treatment with an inhibitor of PERK, GSK2606414 (Axten et al., 2012) (Fig. 6E,F). As above, we confirmed inhibitor activity and specificity by monitoring phosphorylated EIF2α and DDIT3 levels (Fig. 6G,H). Strikingly, treatment with GSK2606414 rescued the reduction in FGFR1 levels induced by hypoxia. This rescue of FGFR1 levels did not result in sustained phosphorylated ERK1/2. Three FGFR1 bands, with different degrees of glycosylation, were detected by immunoblotting (Fig. 6I). The highest molecular weight band (140 kDa), resistant to endoglycosidase H (EndoH) digestion, represents the mature form of FGFR1 (Fig. S5D) (Koika et al., 2013). Although total FGFR1 levels were rescued by GSK2606414 treatment (Fig. 6E), levels of the mature form of the receptor increased to a lesser extent than the immature forms (Fig. 6I-K). The abnormal buildup of the immature forms suggests that receptor maturation is disrupted by hypoxia, providing an explanation for why phosphorylated ERK1/2 levels were not rescued (Fig. 6F).

We next sought to confirm that FGFR1 levels could be rescued by inhibiting PERK in embryos exposed to hypoxia (Fig. 6L-O). We treated pregnant C57BL/6J mice on E9.5 with a single dose of GSK2606414 by oral gavage 1 h before exposure to 5.5% oxygen for the indicated times. As before, low oxygen exposure without GSK2606414 treatment resulted in a significant loss of both total and Golgi-localized FGFR1 protein and, importantly, treatment with GSK2606414 rescued the hypoxia-induced reduction in FGFR1 in the SHF (Fig. 6T,U). Consistent with our in vitro findings, phosphorylated ERK1/2 levels were not sustained in the SHF despite normal FGFR1 levels (Fig. 6P-S,V).

These data indicate that inhibition of FGFR1 protein levels in response to hypoxia occurs specifically by PERK-induced inhibition of translation, both in vitro and in vivo. PERK inhibition restored FGFR1 levels generally, but not the mature active form of FGFR1 and, consequently, not signaling via phosphorylated ERK1/2.

**DISCUSSION**

Almost 200 years ago, Étienne Geoffroy Saint Hilaire showed that gestational hypoxia disrupts embryonic development (Geoffroy
Fig. 6. See next page for legend.
Fig. 6. UPR-mediated translation inhibition mediates FGFR1 loss in response to hypoxic exposure in vitro and in vivo. (A-D) Quantification of expression levels of FGFR1, FLAG-XBP1ΔBD, phospho-EIF2α and Ddit3 in C2C12 treated with 20 μM 4μC8, an inhibitor of Ire1. Scale bar in S: 40 μm in L-O; 80 μm in P-S. (E-H) Quantification of the expression levels of FGFR1, phospho-ERK1/2, phospho-EIF2α and Ddit3 in C2C12 treated with 1 μM GSK2606414, an inhibitor of PERK. (I-K) The PERK inhibitor predominantly rescued the immature forms of the FGFR1. (I) Immunoblot of endogenous FGFR1 in C2C12 cells exposed to hypoxia (0.1% oxygen) and treated with 1 μM GSK2606414 (GSK). Three FGFR1 bands are detected, representing fully glycosylated (140 kDa) and partially glycosylated (120 kDa and 95 kDa) polypeptides. For clarity, irrelevant lanes were removed (vertical lines). (J) Quantification of the absolute levels of mature (circles) and immature (squares) forms of FGFR1. (K) Quantification of the mature, fully glycosylated FGFR1 polypeptide as a percentage of total FGFR1. (L-O) Comparison of FGFR1 (green) protein levels in unexposed and untreated control (L, n=17), unexposed and GSK2606414-treated (M, n=11), hypoxia-exposed and untreated (N, n=16) and hypoxia-exposed and GSK2606414-treated (O, n=17) embryos by immunohistochemistry on sagittal paraffin sections for FGFR1 (green) and GM130 (Golgi, red); nuclei were stained with TO-PRO-3 (red). (P-S) Comparison of phospho-ERK1/2 (green) protein levels in unexposed and untreated control (P, n=16), unexposed and GSK2606414-treated (Q, n=9), hypoxia-exposed and untreated (R, n=16) and hypoxia-exposed and GSK2606414-treated (S, n=15) embryos by immunohistochemistry on sagittal paraffin sections for phospho-ERK1/2 (green); nuclei were stained with TO-PRO-3 (red). (T) Quantification of FGFR1 protein levels in the Golgi of the SHF. (U) Quantification of total FGFR1 protein levels in the SHF. (V) Quantification of phospho-ERK1/2 levels in the SHF. Red dots indicate values for the embryos shown in L-S. **P<0.01, ***P<0.001, ****P<0.0001; ns, not significant. Location of the SHF is indicated by brackets. Scale bar in S: 40 μm in L-O; 80 μm in P-S.

Saint Hilaire, 1820). The cellular and molecular consequences of this finding are only now being elucidated. We have previously shown that short-term gestational hypoxia disrupts FGF signaling in the PSM, altering somitogenesis and leading to vertebral defects (Sparrow et al., 2012). Here we extend this study by demonstrating that short-term gestational hypoxia causes abnormal heart development, resulting in heart defects that are the most common subtype of human CHD. Interestingly, we observed an almost bimodal response to hypoxia, with only ~50% of embryos developing heart defects after hypoxic exposure. This supports the notion that the cause of some of the variation in the clinical outcome of human cases of CHD is environmental in nature. The defects induced by hypoxia are consistent with impaired function of a progenitor population of the heart, namely the SHF. Accordingly, we show that hypoxia reduces FGFR1 protein levels and FGF signaling in the SHF, and also activates the UPR. In both cultured cells and in vivo, we identify that the reduced levels of FGFR1 are likely to be caused by one specific function of the UPR: inhibition of cap-dependent translation. However, it should be noted that we were not able to test whether the third UPR pathway (mediated by ATF6) is also involved. We suggest a model in which the hypoxia-induced UPR results in reduced FGF signaling in the SHF, disrupting heart development and leading to morphological heart defects. We propose that other stressors that trigger the UPR during embryogenesis, such as increased reactive oxygen species or hyperthermia, may also cause CHD. In addition, UPR activation by such stressors may also occur in sites in addition to the SHF. This suggests that activation of the UPR might be a more general disruptor of embryogenesis and thus causes a range of birth defects.

Cells respond to stresses in a variety of ways, of which the UPR is only one. It is formally possible that other cellular stress responses are activated by hypoxia, such as the mTOR pathway (Wouters and Koritzinsky, 2008) and the nucleolar stress response (Boulon et al., 2010; James et al., 2014). The effects of these pathways on heart development could be assessed in future studies.

The hypoxia-mediated induction of the UPR is reversible in our mouse model. This is evident from the reiterative nature of somitogenesis, where each somite is independently formed temporally and spatially. Thus, somites formed after gestational hypoxia ceases are normal, and these give rise to normal vertebrae (Sparrow et al., 2012). By contrast, transient activation of the UPR has lasting morphological effects on the heart. This is because the final structure of the heart relies on many interdependent processes during embryogenesis, with perturbation at any point disrupting its final structure. Thus, although the UPR is reversible during embryogenesis, the effects on organ morphology can be enduring.

One of the primary UPR responses is PERK-dependent attenuation of general protein synthesis. Therefore, it might be expected that a short-term translation block would simply cause a temporary reduction in embryo growth rate. However, the translation block appears to particularly affect the levels of some proteins but not others. In doing so, it disrupts specific processes and leads to distinct morphological defects. One explanation for this is that the UPR-induced translation block specifically affects cellular processes that are reliant on a rapid replenishment of proteins. For example, FGFR1-mediated signaling results in receptor degradation, and for signaling to continue more FGFR1 needs to be translated (Haugsten et al., 2005). Thus, FGFR1 cell surface replacement would be prevented by PERK-induced translation inhibition. This is in contrast to FGFR4, where receptor recycling, rather than degradation, occurs after activation. We might expect receptors of this type to be unaffected by translation inhibition. Indeed, this might explain why expression levels of PDGFRβ protein, another receptor tyrosine kinase expressed in the SHF, like FGFR1, were not affected by hypoxia within the experimental time frame. This differential effect of the UPR on receptor expression and function is an important area for future investigation.

We show that the hypoxia-induced reduction of FGFR1 is due to activation of the UPR and the PERK-dependent inhibition of translation. Inhibition of PERK rescued FGFR1 in cultured cells and in vivo, but did not rescue the reduction of phosphorylated ERK1/2. This can be explained by the fact that, although total FGFR1 levels are restored, the mature form of the receptor is significantly underrepresented. This is not surprising, as although GSK2606414 relieves the inhibition of translation, hypoxia would still inhibit protein folding and maturation in the ER and Golgi (Koritzinsky et al., 2013).

Potentially any stressor present during gestation that triggers the UPR in a tissue where FGFR1 is active could disrupt FGF-dependent developmental processes and lead to birth defects. Accordingly, our studies show that gestational hypoxia at E9.5 in mouse disrupts FGF signaling in the precursor tissues of the heart and vertebrae leading to defects in these structures (this study; Sparrow et al., 2012). The development of other organs might well be disrupted, as FGF/FGFR1 signaling has many key roles in embryogenesis (Thissen and Thissen, 2005; Itoh and Ornitz, 2011). More broadly, our findings suggest that any gestational stress that activates the UPR could disrupt any number of developmental processes and lead to birth defects. The types and numbers of defects that arise will be affected by the timing, duration and degree of the stress during embryogenesis, and the robustness of the developmental processes to withstand perturbation. In humans, different types of birth defects can arise with a non-random co-occurrence. For example, patients with a spectrum of various
combinations of ‘vertebral anomalies, anal atresia, cardiac defects, tracheoesophageal fistula and/or esophageal atresia, renal and radial anomalies and limb defects’ may be classified as having VACTERL association (OMIM 192350). It is possible that the critical stages of development of these organs are coincident. If so, activation of the UPR during this stage of embryogenesis might result in VACTERL.

How likely is it that the UPR is triggered during gestation? In addition to hypoxia, the UPR is activated in response to a variety of environmental stressors such as pollution and pathological conditions including maternal diabetes, viral infection and hyperthermia (Kitamura, 2013). Strikingly, many of these are identified as environmental risk factors for causing congenital malformation in humans (Jenkins et al., 2007). For example, offspring of mothers with pre-gestational diabetes have a significantly increased risk of CHD, with a relative risk of 3-18 (Jenkins et al., 2007), and the embryos of diabetic mice show significant activation of the UPR (Zhao, 2012). In addition, the stressors hyperthermia and hypoxemia are both associated with increased risk of birth defects in humans (Jenkins et al., 2007) and with activation of the UPR in cell culture (Kitamura, 2013). Thus, our results might provide a mechanism for the induction of birth defects by a wide array of environmental factors.

Finally, our in vitro system could be used as the basis of a high-throughput screening strategy to identify potential prophylactic agents to reduce the risks of CHD following the folate/neural tube defect paradigm (MRC Vitamin Study Research Group, 1991). Such prophylactic agents could then be validated in our in vivo system. This has an encouraging precedent, as oral treatment targeting the UPR abrogates the development of clinical prion disease in mice (Moreno et al., 2013). Such an approach has the potential to significantly reduce the incidence of human congenital defects.

MATERIALS AND METHODS
Animal experiments
All animal experiments were performed in accordance with protocols approved by the Garvan Institute of Medical Research/St Vincent’s Animal Experimentation Ethics Committee, Sydney, Australia (approval 12/33). C57BL6/J dams and sires were mated. Pregnant mice were exposed to reduced oxygen levels at normal atmospheric pressure as described (Sparrow et al., 2012). After exposure, the mice were either sacrificed and embryos harvested, or mice were returned to normoxia for embryo harvest at a later date. For GSK2606414 experiments, pregnant mice were dosed with 50 mg/kg by oral gavage on E9.5, 1 h prior to hypoxic exposure. This dose was previously shown to effectively prevent UPR-mediated translational repression in vivo (Moreno et al., 2013).

Heart morphology
Heart morphology was determined using paraffin sectioning or magnetic resonance imaging (MRI) as previously described (O’Reilly et al., 2014), or by optical projection tomography (OPT) as described in the supplementary Materials and Methods. Heart morphology for all samples was assessed by the same observer, with classification of heart defects confirmed by an independent observer.

TUNEL staining and immunohistochemistry
Paraffin sections were labeled with digoxigenin using the ApopTag Fluorescein In Situ Apoptosis Detection Kit (Abcam). Labeled DNA fragments were detected by anti-digoxigenin-alkaline phosphatase, Fab fragments (Roche Life Sciences) and NBT/BCIP solution (Roche Life Sciences). We performed whole-mount immunofluorescence on fixed embryos and sections as previously described (Geffers et al., 2007; Moreau et al., 2014). For details of immunohistochemistry, including quantitation methods, see the supplementary Materials and Methods and Table S2.

RNA in situ hybridization
We performed RNA whole-mount in situ hybridization as described previously (Dunwoodie et al., 1997) using the probes listed in the supplementary Materials and Methods.

Quantitative RT-PCR
Levels of spliced and unspliced Xbp1 transcripts in RNA extracts from E9.5 whole embryos were determined using custom TaqMan gene expression assays (Applied Biosystems) designed using published sequences (Hayashi et al., 2007). RNA extraction, reverse transcription and qPCR were performed as previously described (O’Reilly et al., 2014).

Cell culture and immunoblot
Mouse muscle satellite C2C12 cells were cultured in low oxygen using an H35 Hypoxstation (Don Whitley Scientific). Culture medium was equilibrated in 0.1% oxygen overnight prior to use. Culture dishes were transferred to the workstation, rinsed once with hypoxia-equilibrated medium, and incubated in fresh hypoxia-equilibrated medium for the indicated times. For the study of UPR inhibitors, cells were incubated for 2 h in normoxic or hypoxia-equilibrated medium containing inhibitor. For detection of Xbp1 splicing in the 4µc experiments, a plasmid encoding FLAG-tagged XBP1ΔDBD-venus (Iwawaki et al., 2004) was transiently transfected into C2C12 cells using Lipofectamine LTX and PLUS reagent (Life Technologies) 24 h prior to hypoxic exposure. For further details, see the supplementary Materials and Methods.

Statistical analyses
All statistical analyses were performed with Prism 6 (GraphPad Software). Gene and protein staining intensity from RNA whole-mount in situ hybridizations, immunohistochemistry and immunoblot experiments were compared using Kruskal–Wallis one-way ANOVA with Dunnett’s post-hoc test to compare all treatment groups with the control, except for the data in Fig. 4L and Fig. 6A-J, which were tested with ANOVA followed by Tukey’s post-hoc test to compare the means of each group with every other group. The linear correlation between distal OFT length and the inner angle between proximal and distal OFT (Fig. 1) was tested using non-parametric Spearman correlation. The statistical significance of the prevalence of heart defects or resorbed embryos in exposed versus unexposed embryos (Table 1) was tested using a one-tailed Fisher’s exact test. In figures, data are presented as mean±s.d.

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Competing interests
The authors declare no competing or financial interests.

Author contributions

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References


Feldman, D. E., Chauhan, V. and Koong, A. C. (2002). And...


