Murine FGFR-1 is required for early postimplantation growth and axial organization

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We have explored the role of fibroblast growth factor receptor 1 (FGFR-1) in early embryonic development using three experimental systems: genetically deficient mice, in vitro blastocyst culture, and FGFR-1-deficient embryonic stem cells. Using these systems, we demonstrate that FGFR-1 is required for proper embryonic cell proliferation and for the correct axial organization of early postimplantation embryos but not for mesoderm formation. FGFR-1-deficient embryos display severe growth retardation both in vitro and in vivo and die prior to or during gastrulation. Although these mutants can form nonaxial tissues, such as the allantois, amnion, and yolk sac mesoderm, they display defective patterning of the primitive streak and other axial structures, and frequently exhibit truncations or disorganization of posterior embryonic regions. Such abnormalities are unlikely to be caused by intrinsic blocks in mesodermal differentiation, as FGFR-1-deficient ES cell lines form teratomas consisting of many mesodermal cell types.

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Fibroblast growth factors (FGFs) constitute a family of at least nine structurally and functionally related growth factors that have been implicated in a variety of important biological processes such as mitogenesis, angiogenesis, cell migration, differentiation, and mesoderm induction (Burgess and Maciag 1989; Klagesbrun 1989, Basilico and Moscatelli 1992). High-affinity receptors (FGFRs) for FGFs comprise a family of at least four transmembrane proteins with intrinsic tyrosine kinase activity (Basilico and Moscatelli 1992, Johnson and Williams 1993). These receptors share common structural features, including a hydrophobic leader sequence, three immunoglobulin-like domains, an acidic region, a transmembrane region, and divided tyrosine kinase domains. Numerous mRNA isoforms of the fgr genes are generated by alternative splicing in the extracellular, juxtamembrane, and intracellular domains (Johnson et al. 1990, Hou et al. 1991, Johnson et al. 1991, Werner et al. 1992, Xu et al. 1992).

In the mouse, the mRNA expression patterns of FGFs and FGFRs have suggested potential roles for these genes in early embryogenesis. Several studies have demonstrated that int-2 (FGF-3) is expressed in early migrating mesoderm and neuroectoderm (Wilkinson et al. 1986, Niswander and Martin 1992), that FGF-4 is expressed in the inner cell mass and early mesoderm (Hébert et al. 1990, Niswander and Martin 1992), and that FGF-5 is expressed in the primitive ectoderm [Haub and Goldfarb 1991, Hébert et al. 1991]. Similarly, in situ hybridization studies have shown that fgr-1 and fgr-2 are expressed in gastrulation stage embryos [Orr-Utterger et al. 1991, Yamauchi et al. 1992], that fgr-3 is expressed during organogenesis and limb formation [Peters et al. 1993], and that fgr-4 is expressed in the definitive endoderm and the somitic myotome [Stark et al. 1991].

Studies of Xenopus embryogenesis have demonstrated an important role for the FGF signaling pathway in mesoderm formation. In particular, several members of the FGF family have been shown in animal cap assays to induce ventral mesoderm, such as muscle, mesenchyme, and blood [Kimelman and Kirschner 1987, Paterno et al. 1988, Slack et al. 1989, Smith 1989]. Conversely, expression of an injected dominant-negative FGFR construct frequently inhibits the formation of ventral and posterior mesoderm in intact embryos [Amaya et al. 1991]. Detailed analysis of the effects of this construct in embryos and in animal caps suggests that FGF signaling is essential for many early molecular responses to mesoderm formation.
induction in *Xenopus* [Amaya et al. 1993; Cornell and Kimelman 1994; LaBonne and Whitman 1994].

These findings indicate that the FGF signaling pathway is essential for normal embryonic pattern formation. However, such studies have not yet provided a detailed understanding of the roles of individual FGFs and FGFRs in vertebrate development. One approach for investigating this issue is to mutate individual *fgf* and *fgfr* loci through targeted gene disruption in mice. For example, targeted mutation of *int-2* has revealed essential roles for this gene in inner ear and tail development [Mansour et al. 1993].

As described below, we have created mice that carry germ-line mutations of the *fgfr-1* locus through homologous recombination in embryonic stem (ES) cells. The *fgfr-1* gene was mutated by the insertion of a *neo* gene into its fourth exon, which is shared by all major splice isoforms [Johnson et al. 1991], thereby creating a strong reduction or loss-of-function mutation. The consequences of this mutation are severe, resulting in homozygous lethality before or during gastrulation. Our data demonstrate that signals mediated by FGFR-1 are essential for cell proliferation and axial organization of embryos during postimplantation development.

**Results**

**Targeted disruption of the *fgfr-1* gene**

To create the *fgfr-1* mutation, we generated the targeting construct pFGFR1neo [Fig. 1]. This construct contains a 9-kb *fgfr-1* genomic sequence interrupted in exon 4 by a *PGKNeo* cassette. The fourth exon contains part of the Ig2 domain and part of the hinge connecting the Ig2 and Ig3 domains, which are shared by all major *fgfr-1* splice isoforms [Johnson et al. 1990, 1991; Werner et al. 1992]. The *neo* gene was placed in the opposite orientation relative to that of *fgfr-1* to introduce stop codons in all three reading frames and to prevent potential readthrough transcription from the *PGK* promoter. Therefore, the *PGKNeo* insertion should truncate the *fgfr-1*-coding sequence immediately after the Ig2 domain and should disrupt all major isoforms of the gene.

Three ES clones carrying the targeted *fgfr-1* mutation were injected into C57Bl/6J blastocysts to obtain germ-line transmission. Injection of one of these clones, 11-22, generated 12 high-degree chimeras, with 10 males and 2 females. When mated to black female mice (NIH Black Swiss), nine of the male chimeras gave agouti

![Figure 1. Targeted disruption of the *fgfr-1* gene.](image-url)

[A] Targeting vector pFGFR1neo contains a 9-kb genomic sequence of the *fgfr-1* genomic locus [not drawn to scale]. The solid boxes represent *fgfr-1* exons, numbered according to the published map [Johnson et al. 1991]. The crosshatched box represents *PGKNeo* inserted at an *AatII* site in the fourth exon of *fgfr-1*, such that 7 kb of 5′ and 2 kb of 3′ sequence flank the insert. The *fgfr-1* sequences are flanked by copies of the *HSV-TK* genes (stippled boxes) and the *pUC* vector (open box). Transcriptional directions of *neo* and TK are indicated by arrows. The targeting vector was linearized at the unique *SalI* site prior to electroporation. Homologous recombination within *fgfr-1* would replace the endogenous exon 4 with *PGKNeo*, creating a bandshift from 9 to 6 kb upon EcoRV digestion. The DNA probe used for Southern blotting is a 2-kb *XhoI-EcoRV* fragment, indicated by a solid bar. [B] Targeting vector pFGFR1hyg contains a 6-kb *EcoRI* fragment of *fgfr-1*. A *hyg* gene is inserted at the *AatII* site in the fourth exon of *fgfr-1*. pFGFR1hyg was linearized at the *SalI* site and transfected into 11-22, an ES cell line heterozygous for a *Hyg* allele. *Hyg* was integrated into the *fgfr-1* genomic locus through homologous recombination in embryonic stem (ES) cells. The *fgfr-1* gene was mutated by the insertion of a *neo* gene into its fourth exon, which is shared by all major splice isoforms [Johnson et al. 1991], thereby creating a strong reduction or loss-of-function mutation. The consequences of this mutation are severe, resulting in homozygous lethality before or during gastrulation. Our data demonstrate that signals mediated by FGFR-1 are essential for cell proliferation and axial organization of embryos during postimplantation development.

![Figure 1](image-url)
FGFR-1-deficient mice

pups. Southern blot analysis indicated that ~50% of agouti offspring had received the \(fgfr-1^{ex4}\) mutation.

**The \(fgfr-1^{ex4}\) mutation results in recessive embryonic lethality**

Mice heterozygous for the targeted mutation \(\text{[}fgfr-1^{ex4}/+\text{]}\) were developmentally and anatomically normal. Twelve of these mice were closely monitored for up to 1 year and were completely normal in terms of growth rate, morphology, health, and fertility (data not shown). When we intercrossed heterozygotes, we found no homozygous \(fgfr-1^{ex4}\) mice among 61 newborns analyzed by Southern blotting (Table 1). In subsequent experiments, we have analyzed >500 progeny of heterozygote intercrosses without detecting any homozygotes (data not shown). These results indicate that the \(fgfr-1^{ex4}\) mutation is a recessive embryonic lethal.

To determine the timing of \(fgfr-1^{ex4}\) homozygous lethality, we dissected embryos from heterozygote intercrosses at different gestational days. From E10.5 to E12.5 (i.e., 10.5 to 12.5 days of gestation), 44 of 61 decidual analyzed contained morphologically normal embryos, whereas the remaining 17 decidual were much smaller in size and contained nearly or completely resorbed embryos (Table 1). In contrast, between E6.5 and E9.5, abnormal embryos were present in some of the smaller decidua, whereas other decidua contained resorptions. After dissecting the abnormal embryos free of maternal tissue, we determined their genotype by polymerase chain reaction (PCR) or Southern blot analysis of DNA extracted from whole embryos or visceral yolk sacs. All normal embryos genotyped were either +/+ or \(fgfr-1^{ex4}/+\) at a 1:2 ratio, and 92% (49/53) of the abnormal embryos were homozygous for the \(fgfr-1^{ex4}\) mutation (Table 1). The remaining abnormal embryos (8%) might have been either natural occurrences in this strain background (Table 2) or incorrectly genotyped by PCR because of maternal contamination of the sample. Thus, these data indicate that loss of \(fgfr-1\) function results in embryonic lethality during early postimplantation development.

**Growth defects observed by in vitro blastocyst culture**

To determine whether preimplantation stages were affected by the targeted mutation, blastocysts were isolated from matings between \(fgfr-1^{ex4}/+\) mice and were genotyped by PCR either before \((n = 10)\) or after in vitro culture \((n = 50)\). Wild type, heterozygous, and homozygous blastocysts were recovered in the expected 1:2:1 ratio (Table 1) and appeared morphologically indistinguishable before culture (Fig. 2A,D). These data indicate that embryos homozygous for the targeted mutation are morphologically normal prior to implantation.

However, we observed a dramatic growth disadvantage of homozygous embryos after they were cultured for several days in vitro. During the first 4 days of culture, there were no obvious differences in the morphology of homozygous, heterozygous, and wild-type blastocysts \((n = 27)\). In this period, cultured blastocysts of all three genotypes formed structures that resembled early egg cylinders. After 5–6 days in culture, however, we observed that the \(fgfr-1^{ex4}\) homozygous blastocyst outgrowths were consistently smaller than wild-type or heterozygous outgrowths. Specifically, five out of seven \(fgfr-1^{ex4}\) homozygous outgrowths (Fig. 2E) were much smaller than the majority of wild-type or heterozygous outgrowths (13/20; Fig. 2B,C), whereas the other two \(fgfr-1^{ex4}\) homozygous outgrowths (Fig. 2F) were similar to the remaining seven wild-type or heterozygous outgrowths. Similar observations were made for a second set of embryos \((n = 23)\;\text{(data not shown)}\). When combined with our in vivo observations of growth retardation in homozygous embryos, as described below, these data are consistent with an early embryonic role for FGFR-1 in stimulating mitogenesis.

### Table 1. Morphologies and genotypes of embryos from crosses between \(fgfr-1^{ex4}/+\) animals

| Age            | Number of embryos/litter | Normal | Abnormal | Resorption | Genotype |
|----------------|--------------------------|--------|----------|------------|----------|
|                |                          |        |          |            | +/+      | +/-     | -/-     |
| E3.5<sup>a</sup>                  | 60                      | 60 (60)| 0        | 0          | 15       | 31      | 14      |
| E5.75-6.0<sup>d</sup>              | 47/6                    | 45     | 2        | 0          | 0        |         |         |
| E6.5<sup>a</sup>                   | 95/11                   | 81     | 12       | 2          | 2        |         |         |
| E7.5                       | 229/25                 | 171 (21)| 32 (6)   | 26         | 6        | 15      | 6       |
| E8.5                       | 258/27                 | 196 (51)| 25 (11)  | 37         | 19       | 33      | 10      |
| E9.5                       | 423/30                 | 297 (30)| 49 (36)  | 77         | 10       | 23      | 33      |
| E10.5–12.5                 | 61/7                    | 44 (20)| 17       | 6          | 14       | 0       |         |
| Total (E6.5–12.5)          | 1066/100               | 789    | 118      | 159        | 21       | 40      | 0       |

<sup>a</sup> Parentheses indicate the number of embryos that have been genotyped.

<sup>b</sup> Resorptions were not genotyped because of difficulty in separating embryonic from maternal tissues.

<sup>c</sup> Blastocysts were genotyped either before \((n = 10)\) or after in vitro culture \((n = 50)\).

<sup>d</sup> Embryos were analyzed by sectioning within whole decidua.
Table 2. Morphologies of embryos generated by mating between wild-type females and heterozygous males

| Age     | Number of litters | Number of embryos | Normal embryos | Abnormal embryos | Resorptions |
|---------|-------------------|-------------------|----------------|------------------|-------------|
| E6.0    | 1                 | 7                 | 7              | 0                | 0           |
| E6.5    | 3                 | 30                | 30             | 0                | 0           |
| E7.5    | 8                 | 85                | 81             | 2                | 2           |
| E8.5    | 2                 | 24                | 19             | 2                | 3           |
| E9.5-E13.5 | 10            | 97                | 96             | 0                | 1           |
| Total   | 24                | 243               | 233            | 4                | 6           |

Developmental delays and resorptions in \( \text{fgfr}^1\text{ex4} \) embryos are apparent by the onset of gastrulation

To examine the histology of egg cylinder-stage embryos, we fixed and sectioned intact decidual swellings of litters between E5.5 and E6.5 (Table 1). When we analyzed six litters from \( \text{fgfr}^{-1\text{ex4}}/+ \) intercrosses at \( E5.75 \) and E6.0, we found that 45 of 47 embryos appeared morphologically normal. The remaining two embryos were delayed more extensively (not shown), either as a consequence of the targeted mutation or the background rate of resorption (Table 2).

By E6.5, however, we found that 15% of the decidua (14/95) contained abnormal or partially resorbed embryos (Table 1). The abnormal embryos displayed significant developmental retardation, with shorter egg cylinders and smaller proamniotic cavities than their normal littermates (Fig. 3). Whereas most normal E6.5 embryos had begun to gastrulate, forming a primitive streak and posterior amniotic fold (Fig. 3C), the developmentally delayed embryos had not yet begun to gastrulate (Fig. 3D, E). Although we have been unable to genotype these embryos because of their size, it is likely that they are \( \text{fgfr}^{-1\text{ex4}} \) homozygotes, because growth retardation was not observed in control crosses between wild-type females and \( \text{fgfr}^{-1\text{ex4}}/+ \) males (Table 2). These observations suggest that a significant portion of \( \text{fgfr}^{-1\text{ex4}} \) embryos display a growth defect by the onset of gastrulation.

From E6.5 onward, a high frequency of resorption in the litters of heterozygous crosses was readily observed (Table 1). Most resorption sites found in earlier stages of gastrulation (E6.5 and E7.5) contained partially resorbed embryos (Fig. 3A), whereas >50% of later resorption sites (E8.5 and E9.5) contained completely resorbed embryos (Figure 3B). Because little or no embryonic tissue could be recovered from these resorption sites, we could not genotype them reliably by Southern blotting or by PCR. However, the sum of abnormal embryos and resorptions from E6.5 to E12.5 (\( n = 277 \)) was approximately one-quarter of the total embryos analyzed at these stages (\( n = 1066 \)), indicating that the resorptions represent one of the phenotypes of FGFR-1 deficiency. As a control for the effect of the mixed genetic background, we determined that the rate of abnormal embryos plus resorptions was 4% in crosses between \( \text{fgfr}^{-1\text{ex4}}/+ \) males and wild-type female littermates (Table 2). These results suggest that the vast majority of resorptions found in matings between \( \text{fgfr}^{-1\text{ex4}}/+ \) mice represent homozygous mutant embryos.

Phenotypic variability during gastrulation

Although a significant number of presumptive \( \text{fgfr}^{-1\text{ex4}} \)
homozygotes were resorbed before gastrulation, many mutant embryos survived after E6.5 and were recovered as late as E9.5. At E7.5, most normal embryos (n = 171) were at the late primitive streak stage and some had advanced further (Fig. 3F). In contrast, all abnormal embryos (n = 32) were growth retarded, with some displaying an unusually prominent invagination at the circumferential constriction that divides the egg cylinder between embryonic and extraembryonic ectoderm (Fig. 3G,H).

At E8.5, the ratio of resorptions to abnormal embryos increased, indicating that more fgfr-1<sup>+/−</sup> homozygotes were dying (Table 1). Some of these mutants were extremely small, resembling those found in E7.5 litters (Fig. 4B). Other homozygotes had an extraembryonic portion that was significantly larger than their embryonic portion, so that they resembled a “comma” (Fig. 4A). These mutants displayed little identifiable mesodermal differentiation and represented the majority of abnormal embryos found at E8.5. Histological sections demonstrated that these embryos had abnormally thickened primitive streak regions, with limited outgrowth of extraembryonic mesoderm (Fig. 5B,C).

Significant phenotypic diversity was observed in fgfr-1<sup>+/−</sup> homozygotes isolated from E9.5 litters. Approximately two-thirds of mutant embryos seemed to be derived from the growth of comma-shaped embryos, having formed an abnormal visceral yolk sac that was significantly larger than the embryonic portion (Fig. 4E). These embryos often appeared to have very little poste-
Nonaxial mesoderm differentiation in homozygous embryos

In general, we found that homozygous fgfr-1ex4 embryos were capable of forming nonaxial mesodermal tissues such as allantois, amnion, visceral yolk sac mesoderm, and blood. The allantois of fgfr-1ex4 homozygotes was usually well-formed, although sometimes exaggerated in size [Fig. 5E,F,H]. In contrast, we observed that the visceral yolk sac of fgfr-1ex4 homozygotes was always significantly smaller and abnormal. The surface of the visceral yolk sac of fgfr-1ex4 homozygotes was noticeably bumpy and formed characteristic “ruffled” structures. This visceral yolk sac ruffling may be a consequence of underproliferation of the inner layer of extraembryonic mesoderm with respect to the outer layer of visceral endoderm [Fig. 5H]. Despite the limited growth of yolk sac mesoderm, however, some E9.5 fgfr-1ex4 homozygotes formed blood islands and circulating nucleated erythrocytes [Fig. 5I]. One of the most advanced fgfr-1ex4 homozygotes possessed an abnormal heart with internal blood cells [Fig. 5G].

In addition to formation of nonaxial mesodermal tissues, fgfr-1ex4 homozygotes also displayed some morphological differentiation of the head. Thus, by E9.5, the most advanced fgfr-1ex4 homozygotes possessed structures that resemble head folds prior to anterior neuropore closure [data not shown]. These headfolds were never normal in size or morphology [Fig. 5H], compared with normal embryos [Fig. 5A,D]. Other anterior structures that have been occasionally observed include apparent otocysts and branchial arches [not shown].

**Disorganized axial structures in homozygous embryos**

In contrast to the relative ability of fgfr-1ex4 homozygotes to form head structures and nonaxial mesodermal tissues, we observed significant defects in their ability to differentiate and organize axial and paraxial structures of the trunk and tail [Figs. 4C and F; 5F and G]. Even in the most advanced fgfr-1ex4 embryos that have an apparent anterior–posterior axis, the posterior trunk regions were frequently disorganized and displayed folding of ectodermal layers, again suggesting mismatched growth rates of cell layers [Fig. 5G,H]. Most notably, we have never observed proper differentiation of a neural tube, formation of overt somites, or signs of limb bud formation.

To further characterize these defects in axial organization, we employed whole-mount in situ hybridization to examine two markers for midline structures. First, we used a riboprobe from the Brachyury gene to examine primitive streak and notochord differentiation in fgfr-1ex4 homozygotes [n = 8]. At E7.5, Brachyury is normally expressed in the primitive streak, node, and head process, and at E8.5 and E9.5 it is observed in the tail bud and notochord [Wilkinson et al. 1990; Herrmann 1991] [Fig. 6A [left], B,C]. In fgfr-1ex4 homozygotes, only a patch of staining was evident at E7.5 [Fig. 6A, right], whereas a thicker band of staining was commonly found at E8.5 [Fig. 6D], consistent with the abnormally thickened primitive streak region observed in histological sections [Fig. 5C]. At E9.5, fgfr-1ex4 homozygotes that lacked overt axial organization [flattened disks] often displayed large patches of disorganized Brachyury expression [Fig. 6E]. Homozygotes possessing an overt anterior–posterior axis often showed an excess of midline Brachyury staining [Fig. 6F], presumably corresponding to streak- or notochord-like structures.

These defects in midline structures were confirmed by using a riboprobe from the Sonic hedgehog (Shh) gene, which is normally expressed in the notochord, floor plate, foregut, hindgut, and ventral diencephalon at E8.5 and E9.5 [Fig. 6G,H; Echelard et al. 1993]. At E9.5, fgfr-1ex4 homozygotes [n = 5] displayed profound abnormalities in presumptive notochordal staining [Fig. 6J [arrowhead]], again demonstrating defective axial organization.
Figure 5. Histological sections of embryos at E8.5 and E9.5. [A–C] E8.5 embryos; [D–H] E9.5 embryos. [A,D] fgfr-1<sup>ex4</sup>/+ heterozygotes; all other panels show fgfr-1<sup>ex4</sup> homozygotes. [I] Nucleated embryonic blood cells [nbc] from the heart of a E9.5 fgfr-<sup>ex4</sup> homozygote. The arrowhead in F indicates an abnormal neural plate. Arrowheads in G and H indicate regions of apparent ectodermal folding. [al] Allantois; [am] amnion; [em] embryonic portion of embryo; [hf] head fold; [hr] heart; [me] mesencephalon; [ps] primitive streak; [so] somites; [te] telencephalon; [ys] yolk sac. Bar, 70 μm in A, D, E, G, and H; 18 μm in B and C; 55 μm in F; 3.5 μm in I.

As with the Brachyury probe, we frequently observed expanded bands of midline staining, suggesting an excess of presumptive axial mesoderm [Fig. 6K,L]. Whereas all homozygous embryos analyzed displayed some Shh expression, we never found structures that resembled normal notochord or floor plate. These observations suggest that fgfr-1<sup>ex4</sup> homozygotes are able to differentiate cells with primitive streak- and notochord-like characteristics, albeit in a severely disorganized fashion.

FGFR-1-deficient ES lines can differentiate normally into mesodermal cell types

One conceivable explanation for the lack of somite formation and other mesodermal defects in fgfr-1<sup>ex4</sup> homozygotes is that these abnormalities are attributable to intrinsic blocks in mesodermal differentiation pathways. To determine whether this is the case, we tested whether ES cells lacking FGFR-1 can differentiate normally into a variety of mesodermal cell types. For this purpose, we have generated ES cell lines with two disrupted fgfr-1 alleles by targeting the wild-type allele in heterozygous ISS cells (fgfr-lex4/+) with the construct pFGFR1hyg (Fig. 1B). The fgfr-1<sup>ex4</sup>/fgfr-1<sup>ex4</sup> ES cell lines generated by this targeting experiment are morphologically indistinguishable and have similar growth rates in vitro compared with the parental fgfr-1<sup>ex4</sup>/+ line (not shown).

We have tested the ability of these ES cell lines to differentiate in vivo by generating teratomas through subcutaneous injection into nude mice. Regardless of their genotypes, fgfr-1<sup>ex4</sup>/fgfr-1<sup>ex4</sup>, fgfr-1<sup>ex4</sup>/+ , and wild-type ES cells formed tumor masses that were visible between 1 and 2 weeks following injection, becoming 1–2 cm in diameter after 3–4 weeks. Histological examination indicated that teratomas formed from these cells consisted of many different cell types, including squamous and glandular epithelium, neurons, and cartilage (data not shown). In particular, the fgfr-1<sup>ex4</sup>/fgfr-1<sup>ex4</sup> ES cells could form muscle cells that are histologically indistinguishable from those formed by the fgfr-1<sup>ex4</sup>/+ ES cells (Fig. 7). We therefore conclude that disruption of fgfr-1 is unlikely to result in a significant intrinsic defect in the differentiation of many mesodermal cell types. Instead, the mesodermal defects found in fgfr-1<sup>ex4</sup> homozygous embryos are probably an indirect consequence of abnormal pattern formation.

Discussion

Distinct requirements for FGFR-1 in early embryonic development

We have observed phenotypic defects in fgfr-1<sup>ex4</sup> homozygotes in many tissues and cell types that are known to express FGFR-1. In wild-type embryos, transcripts for fgfr-1 are first detectable at the blastocyst stage (Campbell et al. 1992), and are then found in the primitive ectoderm of the egg cylinder and in migrating embryonic mesoderm during gastrulation (Orr-Urtreger et al. 1991). Later, starting with somitogenesis, fgfr-1 is actively transcribed in both the mesoderm and neuroectoderm (Ya-
Deng et al.

Figure 6. Whole-mount in situ hybridization of wild-type and fgfr-1~ex4 homozygous embryos. (A–F) Hybridization with a Brachyury riboprobe. (A) Wild-type and fgfr-1~ex4 homozygous E7.5 embryos. (B) Wild-type E8.5 embryo. (C) Wild-type E9.5 embryo [dark field]. (D) fgfr-1~ex4 homozygous E8.5 embryo. (E) fgfr-1~ex4 homozygous E9.5 embryo lacking well-defined anterior–posterior axis, showing large patches of scattered staining. (F) fgfr-1~ex4 homozygous E9.5 embryo with diffuse broad midline staining. (G–L) Hybridization with a Sonic hedgehog riboprobe. (G) Wild-type E8.5 embryo; (H) wild-type E9.5 embryo; (I–J) fgfr-1~ex4 homozygous E9.5 embryos. In J, positive staining is indicated by the arrowhead. In L, note the diffuse midline staining of the presumptive notochord (arrow), which resembles the Brachyury staining in F. The arrowhead in L points to presumptive hindgut staining. (al) Allantois; (di) diencephalon; (hi) hindgut; (nt) notochord; (ps) primitive streak; (tb) tailbud; (ys) yolk sac. Bar, 35 μm in A and G; 30 μm in B, F, J, and L; 65 μm in C and H; 20 μm in D and K; 25 μm in E and I.

maguchi et al. 1992). This expression pattern is consistent with observed defects of fgfr-1~ex4 homozygotes in the proliferation of primitive ectoderm and differentiation of embryonic mesoderm.

Somewhat surprisingly, however, our analysis suggests that FGFR-1 is required for two distinct functions in early embryogenesis. First, the appearance of resections and growth retardation by the onset of gastrulation indicates that the earliest deaths of fgfr-1~ex4 homozygotes are not a consequence of defects in mesoderm differentiation or patterning. Instead, we believe that these early abnormalities result from insufficient responses to mitogenic stimuli, because of disruption of the FGF signaling pathway. Consistent with this view, all fgfr-1~ex4 homozygotes recovered at later embryonic stages display severe developmental delays and growth deficiencies. Notably, we find that the requirement for fgfr-1 in cell proliferation appears greater in embryonic tissues, consistent with the observed expression pattern of the gene (Yamaguchi et al. 1992). Cell proliferation may be most dependent on FGFR-1 in mesodermal tissues, as suggested by the folding or ruffling of nonmesodermal cell layers in the head and visceral yolk sac of fgfr-1~ex4 homozygotes. Furthermore, fgfr-1~ex4 blastocyst outgrowths demonstrate noticeable growth retardation after several days of in vitro culture. This observation strongly argues that the growth abnormalities of fgfr-1~ex4 embryos in

Figure 7. Histological sections of teratomas derived from fgfr-1~ex4/+ and fgfr-1~ex4/fgfr-1~ex4 ES cells. Shown are muscle cells obtained from injection of 11-22, an fgfr-1~ex4/+ ES cell line [A], and from injection of 23-46, an fgfr-1~ex4/fgfr-1~ex4 ES cell line [B]. Bar, 18 μm.
vivo represent intrinsic defects and are unlikely to result from blocks in maternal-embryonic interactions. Our results therefore indicate that FGFR-1 is required for transmitting important mitogenic stimuli in early embryogenesis.

FGFR-1-deficient embryos have extensive abnormalities in both axial and paraxial structures such as the primitive streak, notochord, head folds, and somites, as demonstrated by histological analysis and by whole-mount in situ hybridization. In principle, some of these abnormalities may simply represent a consequence of early growth defects of specific cell types or structures that are essential for axial organization. If this is the case, FGFR-1 might function solely as a mediator of mitogenic stimuli and would not be directly involved in differentiation or pattern formation. However, it is difficult to see how defects in cell proliferation could account for phenotypes such as the accumulation of cells in the primitive streak and the expansion of axial mesoderm.

Alternatively, FGFR-1 may have an additional function distinct from promoting cell proliferation, such as specifying mesodermal differentiation and/or patterning. Because differentiation of FGFR-1-deficient ES cell lines proceeds normally in teratomas, it is unlikely that FGFR-1-deficient cells possess intrinsic blocks in mesodermal differentiation pathways. Instead, defects in the allocation of progenitor cells among alternative pathways of mesoderm differentiation are more consistent with the phenotypes of FGFR-1-deficient embryos. Thus, one of the earliest phenotypes observed was the formation of an abnormal primitive streak region, as shown by the prominent invagination and by the accumulation of cells in the streak (Fig. 5C). It is known that different mesoderm subpopulations are derived from cells migrating from different regions of the primitive streak (Tam 1989; Lawson et al. 1991; Lawson and Pedersen 1992). Abnormalities in the primitive streak region might cause alterations in migration patterns or pathways of primitive cells, which, in turn, change cell fate and severely affect subsequent mesoderm formation and differentiation. For example, the expansion of the presumptive axial mesoderm at the expense of the paraxial mesoderm could be caused by a lateral migration failure of cells at the anterior end of the streak. This would be consistent with the observed increase in midline expression of Brachyury and Shh, and with the reduction of yolk sac mesoderm and failure to form somites. Thus, in the absence of FGFR-1, the proper allocation of mesodermal progenitors among different embryonic regions and/or differentiation pathways may be disrupted. Such defects in cell fate allocation might be analogous to the formation of neuroectoderm in Xenopus embryos blocked in activin receptor signaling (Hemmati-Brivanlou and Melton 1992, 1994).

Phenotypic variation and functional redundancy of FGF receptors

Interestingly, we have observed a significant variation in the phenotype of fgfr-1<sup>ex4</sup> homozygotes, which can be accounted for by several possible explanations. First, this phenotypic variation might arise through functional redundancy of FGFR-1 with other members of the FGF receptor family. In vitro binding studies have demonstrated that each receptor gene can produce protein products that can bind to several FGF ligands, with differing binding specificities generated through alternative splicing (Dionne et al. 1990; Mansukhani et al. 1990; Dell and Williams 1992, Duan et al. 1992, Ornitz and Leder 1992; Vainikka et al. 1992, Werner et al. 1992; Ron et al. 1993). For example, all four fgfr genes can express receptors that bind to aFGF with high affinities while binding bFGF and kFGF with varying affinities. Thus, phenotypic variation may result from lower ligand binding affinities of redundant FGFRs, increasing their sensitivity to threshold signaling effects that may be influenced by epigenetic phenomena.

Second, phenotypic variation may be a consequence of a nonuniform genetic background [129/terSV x NIH Black Swiss], resulting in segregation of potential modifying loci. Our preliminary results show that the resorption rate from crosses between fgfr-1<sup>ex4/+</sup> mice increases slightly (to 5%) in the inbred 129/terSV background. However, abnormal embryos resembling those in Figures 3, 4, and 5 have also resulted from these crosses (C.-X. Deng and M.M. Shen, unpubl.). These findings suggest that a mixed genetic background cannot be completely responsible for the phenotypic variation in fgfr-1<sup>ex4</sup> homozygotes.

Finally, phenotypic variation might be attributable to the expression of a severely truncated gene product. However, the fgfr-1<sup>ex4</sup> mutation severely truncates all known FGFR-1 isoforms, except for a single isoform that only possesses an Ig1 domain (Eisemann et al. 1991), which is poorly characterized. It is also conceivable that any potential truncated FGFR-1 products might act in a dominant-negative fashion, perhaps through competition for ligand binding with FGFR-1 or other FGFRs. However, in vitro assays demonstrate that both the Ig2 and Ig3 domains are required for high-affinity ligand binding (Zimmer et al. 1993), whereas the presence or absence of the Ig1 domain has very little effect (Miki et al. 1992). Therefore, we believe our fgfr-1<sup>ex4</sup> disruption is a strong reduction or loss-of-function mutation and represents a candidate null mutation. This conclusion is supported by studies from the accompanying paper in which Yamaguchi and Rossant (this issue) have generated a mouse strain with a similar phenotype by deleting the transmembrane and a part of the catalytic domain of the fgfr-1 gene.

The FGF signaling pathway and mesoderm formation in mouse and frog

Many experiments on Xenopus embryos have supported a role for the FGF signaling pathway in mesoderm induction. Injection of a dominant-negative FGFR construct into intact embryos frequently results in tadpole-
stage animals displaying nearly complete head differentiation, but lacking trunk and tail structures, indicating that FGF signaling is involved in specifying the differentiation of ventral and posterior mesoderm in *Xenopus* [Amaya et al. 1991, 1993]. Further analysis has demonstrated that dominant-negative FGFRs can inhibit the differentiation of more dorsal mesoderm both in intact *Xenopus* embryos and in activin-treated animal caps. Specifically, dominant-negative FGFR receptors can block the expression of *Brachyury* and other markers of non-prechordal mesoderm, including those for skeletal muscle [Amaya et al. 1993; Cornell and Kimelman 1994; LaBonne and Whitman 1994].

To some extent, the phenotypes of *fgfr-1* homozygotes resemble those of *Xenopus* embryos that have been injected with dominant-negative FGFR constructs. In particular, similar pattern truncations are found in some *fgfr-1* homozygotes in which head fold-like structures are juxtaposed to the allantois. However, unlike these *Xenopus* embryos, *fgfr-1* homozygotes possess significantly abnormal head structures. In addition, *fgfr-1* homozygotes can form many extraembryonic mesodermal cell types, such as allantois, amnion, yolk sac mesoderm, and blood, which are derived from “posterior” primitive streak regions in the fate map of the gastrulation-stage mouse embryo [Lawson et al. 1991].

Most notably, however, FGFR-1 is not absolutely required for mesoderm formation in the mouse. For example, FGFR-1 is not required for *Brachyury* expression, as demonstrated by whole-mount in situ analysis. Murine FGFR-1 is also not required for the differentiation of muscle or other mesodermal cell types, as shown by the analysis of FGFR-1-deficient ES cells. Thus, the failure to form somitic mesoderm in FGFR-1-deficient mice is likely to be a consequence of defective mesodermal pattern formation. The basis for these phenotypic differences between mouse and frog may reflect species differences in receptor redundancy or signaling pathways. Alternatively, the potentially broader action of a dominant-negative construct on the entire FGFR receptor family may differ from the effects of a specific disruption of FGFR-1 function.

Our analysis indicates that FGFR-1 is required during mouse embryogenesis for two essential functions: proper embryonic cell proliferation and pattern formation. Although the patterning abnormalities in FGFR-1-deficient mice might be caused either by a mitogenic defect, or by a defect occurring at earlier stages of the gastrulation process, a dual role for FGFR-1 is consistent with biochemical analyses of an FGFR with a point mutation in its cytoplasmic domain at the autophosphorylation site Tyr-766 [Mohammadi et al. 1992, Peters et al. 1992]. This mutated receptor is defective for phospholipase Cγ signaling and phosphatidylinositol hydrolysis but is still able to mediate mitogenic activity in transfected myoblasts. Interestingly, it has been proposed that a high rate of phosphatidylinositol turnover is involved in specifying ventral mesoderm in frog embryogenesis [Berridge et al. 1989]. It is therefore conceivable that FGFR-1 could utilize distinct signal-transducing domains to mediate cell proliferation and axial patterning in mouse development.

**Materials and methods**

**Targeting vectors**

Recombinant phages containing genomic DNA of *fgfr-1* were isolated from a 129 mouse library (Stratagene) by using a mouse *fgfr-1* cDNA probe. Two overlapping fragments containing exon 4, a 9-kb *Xhol–SalI* fragment (the *SalI* site is from the polylinker region of the AFix II phage vector) and a 6-kb *EcoRI* fragment, were subcloned into pBluescript IKKS+1 [Stratagene]. To make the pFGFR1neo targeting vector, we inserted the PGKneo gene [Tybulewicz et al. 1991] into exon 4 of *fgfr-1* at a unique AatII site in the 9-kb *Xhol–SalI* fragment. Two different thymidine kinase genes, *TK-1* and *TK-2* (Mansour et al. 1988; Thomas and Capecchi 1990), were then placed at the 5' and 3' ends of the genomic DNA, respectively (Fig. 1A). The pFGFR1hyg vector was constructed by inserting a pMC1hyg gene (provided by Dr. Teruhisa Tsuzuki, Kyushu University, Japan) into the fourth exon in the 6-kb *EcoRI* fragment followed by the addition of *TK-1* at the 5' end of the genomic DNA [Fig. 1B]. Both targeting vectors were partially sequenced to confirm the predicted insertions of neo and hyg genes (data not shown).

**Homologous recombination in ES cells**

The culture, electroporation, and selection of J1 cells was carried out as described [Li et al. 1992]. ES cell colonies that were resistant to both G418 and FIAU were analyzed by Southern blotting for homologous recombination events. Genomic DNA from these cell lines and the parental cell line was digested with *EcoRI* and probed with a 2-kb 3'-flanking fragment specific for the *fgfr-1* locus. DNA from the parental ES lines showed a 9-kb fragment diagnostic of the wild-type *fgfr-1* gene, but in targeted cell lines this fragment was shifted down to 6 kb as the result of the introduction of an *EcoRV* site in PGKneo (Fig. 1C, lanes 1, 2). Correct targeting was confirmed by digestion with other enzymes (*BamHI* and *EcoRI*), and by using internal probes (data not shown). Of a total of 60 G418/FIAU colonies analyzed, 19 contained the targeted *fgfr-1* disruption. In absolute terms, the targeting frequency achieved by the pFGFR1neo vector was one event per 2 x 10⁵ electroporated ES cells.

The transfection conditions of pFGFR1hyg were identical to those of pFGFR1neo. After electroporation, ES cells were plated on a layer of hygromycin-resistant feeder cells (provided by Dr. Mario Capecchi, University of Utah, Salt Lake City) and selected with hygromycin B at a concentration of 100 μg/ml. ES cells resistant to both hygromycin and FIAU were analyzed for homologous recombination events by Southern blotting with the same probe used in the first round of targeting. Among 90 hyg−/FIAU− clones analyzed, 2 showed the expected band shift, indicating that the wild-type allele was disrupted (Fig. 1C, lanes 3, 4). Additional analysis using multiple enzyme digestions confirmed that the targeting events in these *fgfr-1* clones were accurate. These clones contained no wild-type *fgfr-1* message as assayed by reverse transcriptase (RT)–PCR, using primers spanning the neo insertion site, whereas transcripts in the parental 11-22 ES cell line were readily detectable (data not shown).

**Generation of germ line chimeras**

We superovulated 4-week-old C57Bl/6J females with 2.5 μ of pregnant mare’s serum (Calbiochem) 2 days prior to mating and...
with 2.5 μ hCG (Organon) on the day of mating with C57Bl/6J males. Blastocysts were collected from pregnant C57Bl/6J females at 3.5 days postcoitum. Approximately 15–18 ES cells were microinjected into each blastocyst. The injected blastocysts were implanted into the uteri of pseudopregnant Swiss Webster (Taconic) foster mothers and allowed to develop to term. Male chimeras (identified by the presence of agouti coat color) were test mated with NIH Black Swiss females (Taconic). These animals were used for testing germ-line transmission because they have larger litters and care for their offspring better than C57Bl/6J females (data not shown). Germ-line transmission was confirmed by agouti coat color in the F1 animals, and all agouti offspring were tested for the presence of the fgfr-1<sup>neo</sup> allele by Southern analysis or PCR.

Genotype analysis

For adults and embryos of 8.5 days of gestation and older, we determined most genotypes by Southern blotting using the flanking probe described above. For embryos younger than E8.5, we determined genotypes by PCR. For PCR analysis, embryos or visceral yolk sacs were incubated in 200 μl of extraction buffer [50 mM TrisCl at pH 8.0, 100 mM EDTA, 1% SDS, 100 μg/ml proteinase K] for 1 hr at 50°C. After digestion, samples were extracted once with an equal volume of phenol/chloroform and once with an equal volume of chloroform/isooamyl alcohol (49:1). DNA was precipitated with two volumes of 100% ethanol, together with 10 μg of yeast tRNA as a carrier, and then dissolved in 20–50 μl of water. Two microliters of each DNA sample was amplified in 50 μl of reaction buffer containing fgfr-1 primer 1 (5’-ATGGATTCTGTGGTGCCTTC-3’) and primer 3 (5’-AACCTGAAAAGGGAAGGGG-3’). This primer pair flanks the PGK<sup>neo</sup> insertion site and amplifies a 261-bp fragment from the wild-type fgfr-1 gene. Theoretically, these primers could also amplify an ~2100-bp fragment from the fgfr-1<sup>neo</sup> allele in heterozygous or homozygous embryos. However, under our PCR conditions, the amplification of this fragment was inefficient and not easily visualized by ethidium staining. DNA was also amplified using neo primer 1 (5’-AGAGGCTATTCG-GCTATGACTG-3’) and primer 2 (5’-TTCGTCCAGATCAT- CCTGATC-3’) to detect the neo gene in the fgfr-1<sup>neo</sup> allele. In this case, a 431-bp fragment is detected in embryos heterozygous or homozygous for the fgfr-1<sup>neo</sup> allele, whereas no signal can be detected in wild-type embryos.

Embryo analysis

Mice heterozygous for the fgfr-1<sup>neo</sup> allele were mated for timed pregnancies, with the day of the vaginal plug considered to be 0.5 day of development (E0.5). At the desired time points, we dissected embryos from decidual for photography and further analysis. For E8.5 and E9.5 embryos, we collected the visceral yolk sac for genotype determination by PCR. For histology studies, embryos were fixed in 4% paraformaldehyde or Bouin’s fixative (Sigma) at 4°C for 1 hr, dehydrated through a graded alcohol series, and then embedded in paraffin. Sections of 6-μm were prepared and stained with Harris hematoxylin and eosin according to standard procedures.

In vitro culture of blastocysts

Conditions for blastocyst culture were similar to those described [Hsu 1979], except that blastocysts were incubated individually in 24-well plates. Under these culture conditions, ~70% of the blastocysts hatched from the zona pellucida and attached to petri dishes within 24 hr, another 25% attached within 48 hr, and the remaining 5% either died or were lost during changes of the culture medium. Blastocyst outgrowths were inspected daily and photographed to monitor their development. After 6 days of culture, DNA was prepared for genotyping by PCR.

Whole-mount in situ hybridization

Embryos were collected and fixed in 4% paraformaldehyde for 8–14 hr at 4°C, followed by dehydration through a methanol series and storage at −20°C until use. Embryos were then treated and hybridized to digoxigenin-labeled riboprobes essentially as described [Riddle et al. 1993]. The Brachyury probe was synthesized from a plasmid containing a 716-bp fragment from the 3’-untranslated region [bp 1048–1764 of the published sequence] [Herrmann et al. 1990], which was PCR amplified and cloned into the PstI and EcoRI sites of Bluescript II KS(+). For riboprobe synthesis, this plasmid was linearized at the polylinker NotI site and transcribed with T3. The Shh probe was provided by Dr. Andrew McMahon [Echelard et al. 1993].

Differentiation of ES cells in vivo

ES cells heterozygous or homozygous for the fgfr-1<sup>neo</sup> mutation were trypsinized and suspended in PBS at a concentration of 10⁷ cells/ml. Cells (10⁴) were subcutaneously injected into 3- to 5-week-old nude mice. The mice were sacrificed 3–4 weeks later, when tumors were 1–2 cm in diameter. The tumors were fixed in Optimal*Fix (American Histology Reagent Company, Inc.) and processed for histological sectioning.

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