Genetic analysis of the Müllerian-inhibiting substance signal transduction pathway in mammalian sexual differentiation

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Müllerian-inhibiting substance (MIS) is a member of the transforming growth factor-β (TGF-β) gene family. MIS expression in males causes the regression of the Müllerian ducts, an essential process in male sexual differentiation. Recently, an MIS type II receptor gene has been isolated that is expressed during embryogenesis in mesenchymal cells adjacent to the Müllerian duct epithelium and in Sertoli and granulosa cells of the fetal and adult, male and female gonads, respectively. MIS receptor mutant males develop as internal pseudohermaphrodites, possessing a complete male reproductive tract and also a uterus and oviducts, a phenocopy of MIS ligand-deficient male mice. They express both MIS mRNA and protein, showing that ligand was present, but target organs were hormone-insensitive. All produce sperm, but the majority were infertile because the presence of their female reproductive organs blocks sperm transfer into females. Focal seminiferous tubule atrophy accompanied by Leydig cell hyperplasia was observed and began as early as 2 months of age. The phenotype of MIS ligand/MIS receptor double mutant males was indistinguishable from those of each single mutant. MIS receptor/α-inhibin double mutant males developed testicular stromal tumors and large fluid-filled uteri that were identical in phenotype to MIS ligand/α-inhibin double mutant males. These studies provide in vivo evidence that MIS is the only ligand of the MIS type II receptor, in contrast to the complexity of other TGF-β gene family signaling pathways.

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males that lack MIS also develop as internal pseudohermaphrodites with uterine and oviductal tissues, a condition known as persistent Müllerian duct syndrome (PMDS) [Guerrier et al. 1989]. In contrast to MIS ligand-deficient mice, PMDS human males often have undescended testes. The mutant phenotypes of the MIS ligand-deficient mice and PMDS humans highlight the highly specific role of MIS in the regression of the Müllerian ducts during male sexual differentiation.

Transgenic mice that ectopically express high levels of human MIS (hMIS) throughout development have mutant phenotypes restricted to sexual differentiation and the reproductive organs [Behringer et al. 1990]. Female hMIS transgenic mice lack a uterus, oviducts, and ovaries. Ovaries initially form, but lose germ cells. Subsequently, the somatic cells of the germ cell-depleted ovary transdifferentiate into tubule-like structures reminiscent of the seminiferous tubules of the male gonad. Most of the hMIS transgenic males developed normally and were fertile. A proportion of male transgenic mice that express very high levels of hMIS are not masculinized [Behringer et al. 1990]. Externally, they appear female, and internally, their testes are undescended and the Wolffian duct system is not differentiated, suggesting a defect in androgen biosynthesis. Subsequent studies showed that male transgenic mice that ectopically express MIS have depressed levels of circulating testosterone, suggesting that excess hMIS can alter Leydig cell function [Lyet et al. 1995]. The tissue-specific MIS-induced mutant phenotypes of transgenic mice with high levels of circulating MIS suggest that the limiting factor in this signal transduction pathway is the MIS receptor.

Thus, the characterization of the MIS receptor is an important step in understanding the MIS signal transduction pathway in mammalian sexual differentiation and reproductive development.

TGF-β family members signal through two membrane-bound serine/threonine kinases [Wrana et al. 1994]. Type II receptors bind ligand when expressed alone but require the presence of the type I receptor for signaling [for review, see Massagué et al. 1994]. Recently, MIS type II receptor genes have been isolated from rat Sertoli cell (Baanrends et al. 1994; Teixeira et al. 1996) and rabbit fetal ovary cDNA (di Clemente et al. 1994) libraries. The human gene has been cloned from a prepubertal testis cDNA library [Imbeaud et al. 1995] and a human genomic library. Sequence comparisons suggest that the rat, rabbit, and human genes are homologs. The expression of the type II receptor gene is localized to the mesenchymal cells adjacent to the Müllerian duct epithelium during embryogenesis and Sertoli cells and granulosa cells in fetal and adult testes and ovaries, respectively [Baanrends et al. 1994, 1995a, b; di Clemente et al. 1994, Teixeira et al. 1996]. In addition, receptor mRNA was detected in the gravid uterus [Teixeira et al. 1996]. MIS-specific binding with a dissociation constant of 2.48–2.55 nm has been demonstrated in COS cells transfected with the human MIS type II receptor cDNA [Imbeaud et al. 1995]. On the basis of its structure, expression pattern, and in vitro binding data, this gene encodes a type II receptor for MIS.

To investigate the role of this type II receptor in the MIS signal transduction pathway of male sexual differentiation, MIS type II receptor mutant mice were gener-
ated by gene targeting in embryonic stem (ES) cells. Analysis of the mutant phenotypes of MIS receptor homozygous mutants, MIS ligand/MIS receptor double mutants, and MIS receptor/α-inhibin double mutants suggests that in vivo MIS is the only ligand of the MIS type II receptor. These findings are in contrast to the complexity observed for other TGF-β family signaling pathways.

Results

Generation of a MIS type II receptor mutant allele in the mouse germ-line

To mutate the MIS type II receptor gene in mouse ES cells, we generated a targeting vector that deletes 4.4 kb of the receptor locus including the first six exons, by replacing them with a neomycin-resistance expression cassette (Fig. 1A). When the vector recombines with the endogenous gene, novel EcoRV and SacI sites are introduced (Fig. 1A). Correctly targeted clones can, therefore, be detected by the presence of an additional 6.6-kb mutant fragment when digested with EcoRI/EcoRV and hybridized with a 5′ probe external to the region of vector homology or by the presence of a 5.3-kb mutant band when digested with SacI and hybridized with a 3′ probe external to the region of vector homology (Fig. 1B). Correct targeting deletes ~2.0 kb of 5′-flanking sequence and the first six exons of the MIS type II receptor locus that encode the translation initiation start codon, the signal peptide, the transmembrane region, and ~20% of the serine/threonine kinase domain. It was predicted that this mutation would functionally inactivate the MIS receptor. Two correctly targeted ES clones successfully contributed to the germ-lines of chimeric mice generated by blastocyst injection. The phenotypes of the MIS receptor mutant mice from two independently derived ES cell clones were identical. The phenotype of the MIS type II receptor mutation, examined on either a C57BL/6 × 129/SvEv mixed genetic background and a 129/SvEv inbred genetic background, was essentially the same.

Male MIS receptor mutants develop as internal pseudohermaphrodites

Both male and female mice heterozygous for the MIS receptor mutation appeared normal and were fertile. Males and females homozygous for the MIS receptor mutation were recovered from matings between heterozygotes at the predicted Mendelian ratios (Fig. 1C). Thus, MIS receptor homozygous mutant mice were viable. Because altered expression of MIS can change the external sexual phenotype of transgenic mice, the sex chromosome genotype of the MIS receptor mutants was assessed by use of a Y chromosome-specific probe (Behringer et al. 1990). In each case, the sexual phenotype of the MIS receptor mutants matched the Y chromosome genotype.

All MIS receptor homozygous mutant females were normal and fertile, as is the case for MIS ligand-deficient females (Behringer et al. 1994). All of the MIS receptor homozygous mutant male mice had testes of normal size that were fully descended into the scrotal sac. Male-specific reproductive tissues derived from the Wolffian ducts had differentiated normally. These MIS receptor mutant males also developed Müllerian duct derivatives, including a uterus, oviducts, and partial vagina that were superimposed upon the male reproductive system (Fig. 2A,B). These gross anatomical abnormalities were identical to those of the reproductive tracts of MIS ligand-deficient male mice (Fig. 2C). Because these male MIS receptor mutant mice had testes and both Wolffian and Müllerian duct-derived tissues, they are internal pseudohermaphrodites.

To confirm that this mutant phenotype was caused by the lack of MIS signal transduction but not by the lack of MIS itself, MIS mRNA and protein were measured in the MIS receptor-deficient mutants. In males, MIS levels are highest before birth in the fetal testes and persist after birth but at lower levels (Hacker et al. 1995). In females, low levels of MIS are produced in the ovary after birth (Münsterberg and Lovell-Badge 1991). Reverse transcription–polymerase chain reaction (RT–PCR) was used to examine the expression of MIS mRNA in embryonic day 13.5 (E13.5) gonads (Fig. 3). As expected, E13.5 females did not express detectable levels of MIS mRNA. In contrast, both heterozygous and homozygous E13.5 MIS receptor-deficient males expressed MIS mRNA. Therefore, at the stage when Müllerian duct regression should be actively occurring in MIS receptor-deficient males, MIS mRNA is present. In addition, serum levels of MIS protein in MIS receptor-deficient mutants and controls at 9
pressed and regulated in a normal manner in both male and female MIS receptor.

These in vivo results demonstrate that the MIS signal transduction pathway is not essential for male germ cell development and that the infertility observed in the MIS receptor homozygous mutant males is not caused by factors intrinsic to the male germ cells. The initial infertility of the mutants is probably caused by the physical blockage of sperm transport into the female reproductive tract caused by the presence of the Müllerian duct derivatives as observed in MIS ligand deficient mice [Behringer et al. 1994]. Older MIS receptor homozygous mutant males [6–11 months old] developed testicular pathology that also contributed to their inability to sire offspring (see below).

In contrast to the fertility defects observed in the MIS receptor-deficient males, all of the MIS receptor homozygous mutant females [n > 20] were fertile. Normal folliculogenesis was observed in the ovaries of the MIS receptor-deficient females, suggesting that signaling through the MIS receptor is not essential for female germ cell development or folliculogenesis.

Fertility of MIS receptor mutant mice

MIS ligand and MIS receptor are expressed in Sertoli cells of the testis and granulosa cells of the postnatal ovary, suggesting that the MIS signal transduction pathway may also regulate gametogenesis. Indeed, exposure of fetal ovaries to MIS in vitro or in vivo leads to germ cell loss [Vigier et al. 1987; Behringer et al. 1990]. Therefore, MIS receptor-deficient mice might be expected to have fertility defects. To test this hypothesis, matings were established between MIS receptor homozygous or heterozygous mutant males and wild-type females. In addition, MIS receptor homozygous or heterozygous mutant females were bred with wild-type males. All of the males heterozygous for the MIS receptor mutation immediately impregnated their mates. However, less than half (7 out of 19) of the MIS receptor homozygous mutant males sired offspring. The infertile MIS receptor homozygous mutant males mated with females and produced vaginal plugs, but no sperm was found in the plugs. In addition, the MIS receptor homozygous mutant males that were fertile sired fewer litters compared with controls, suggesting that even though they were fertile, their ability to sire offspring was compromised.

Histological analysis of the testes of 1.5- to 6-month-old MIS receptor homozygous mutant males that were infertile revealed no consistent differences in spermatogenesis in comparison with the fertile MIS receptor homozygous mutants, heterozygotes, or wild-type males (data not shown). These results suggest that signaling through the MIS receptor is not essential for male germ cell development and that the infertility observed in the MIS receptor homozygous mutant males is not caused by factors intrinsic to the male germ cells. The initial infertility of the mutants is probably caused by the physical blockage of sperm transport into the female reproductive tract caused by the presence of the Müllerian duct derivatives as observed in MIS ligand deficient mice [Behringer et al. 1994]. Older MIS receptor homozygous mutant males [6–11 months old] developed testicular pathology that also contributed to their inability to sire offspring (see below).

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Table 1. Serum levels of MIS in d9 and d10 offspring

| Sex   | MIS [ng/ml ± S.D.] |
|-------|-------------------|
|       | +/+  | +/-  | -/-  |
| Male  | 108.82 ± 34.74 [8]  | 104.93 ± 34.00 [19]  | 115.00 ± 37.65 [13] |
| Female| 8.50 ± 9.00 [11]   | 8.84 ± 8.13 [14]     | 8.25 ± 12.44 [10]   |

*Numbers in parentheses represent number of animals.
Table 2. Pathology of the testes of MIS receptor mutants

| Age (months) | Normal (no. of animals) | Focal germinal epithelium atrophy (no. of animals) |
|--------------|-------------------------|--------------------------------------------------|
| 1.5–4        | 3                       | 3 [2]                                            |
| 5–7          | 3                       | 6 [6]                                            |
| 8–11         | 6                       | 6 [6]                                            |

*Numbers in parentheses represent number of animals with localized Leydig cell hyperplasia.

Two 8- to 10-month-old mice had a single focus of calcification in the seminiferous tubules.

Two 10-month-old mice had significant loss of spermatogenesis and diffuse Leydig cell hyperplasia.

was correspondingly diffuse Leydig cell hyperplasia. None of the MIS receptor heterozygotes displayed any testicular abnormalities.

**MIS ligand/MIS receptor double mutant mice**

The MIS receptor mutation yields a phenotype in male mice that is a phenocopy of the male MIS ligand mutant phenotype, suggesting that for Mullerian duct regression, MIS is the only functional ligand of the MIS type II receptor. To further examine this hypothesis, we generated MIS ligand/MIS receptor double mutant mice. We predicted that if the MIS ligand only interacted with the MIS receptor and that if the receptor only interacted with the MIS ligand, then the elimination of both the MIS ligand and the MIS receptor should result in no novel mutant phenotypes.

MIS ligand/MIS receptor compound heterozygous mutant males and females were normal and fertile. These compound heterozygotes were interbred to generate mice homozygous mutant for both the MIS ligand and MIS receptor mutations. As shown in Figure 2D, the gross anatomy of the reproductive tissues from a MIS ligand/MIS receptor double homozygous mutant male is indistinguishable from those of MIS ligand- or MIS receptor-deficient male mice (Fig. 2B,C). Histological analysis showed no major differences in spermatogenesis in these double homozygous mutant males in comparison to controls (data not shown). The fertility of MIS ligand/MIS receptor double homozygous mutant males was examined by mating with wild-type females. Six out of six MIS ligand/MIS receptor double mutant males were infertile. All of the MIS ligand/MIS receptor double homozygous mutant females [n>10] were fertile and produced litters of normal size at normal intervals. These results clearly show that the mutant phenotypes of the MIS receptor mutation and the MIS ligand/MIS receptor compound mutations are phenocopies of the MIS ligand mutant phenotype. This indicates that MIS is the only ligand for the MIS type II receptor.

**Synergistic action of MIS and inhibin signaling pathways in testicular tumorigenesis**

α-Inhibin is a member of the TGF-β gene family that is expressed in multiple tissues including the somatic cells of the male and female gonads (Vale et al. 1990). Male and female α-inhibin-deficient mice develop gonadal tumors of Sertoli and granulosa cell origin (Matzuk et al. 1992). Because MIS and inhibins are related hormones, are produced by the same gonadal cell types, and both loss-of-function mutants develop gonadal abnormalities, it was postulated that there might be cross talk between the MIS and the inhibin signaling pathways. This hypothesis was examined previously by generating MIS ligand/inhibin double homozygous mutant mice (Matzuk et al. 1995a). MIS ligand/inhibin double homozygous mutant male mice developed testicular tumors and large fluid-filled uteri. Their testes developed Sertoli/granulosa cell tumors and Leydig cell neoplasia earlier than α-inhibin-deficient and MIS ligand-deficient controls, suggesting that MIS and inhibins synergize to influence testicular tumorigenesis (Matzuk et al. 1995a).

To understand the level of interaction between these two signaling pathways, MIS receptor/α-inhibin double-mutant mice were generated.

The most prominent phenotype observed in the MIS receptor/α-inhibin double homozygous mutant males was a dilated fluid-filled uterus and large testicular tumors, that were identical in phenotype to those observed in MIS ligand/α-inhibin double homozygous mutant male mice (Fig. 5A,B). α-Inhibin-deficient mice also develop testicular tumors but the tumors are more hemorrhagic and never as large as the tumors in either MIS ligand/α-inhibin or MIS receptor/α-inhibin double homozygous mutant males [Fig. 5C, D]. Histologically, multifocal Leydig cell neoplasia and multifocal Sertoli/granulosa cell tumors were observed in the testes of MIS receptor/α-inhibin double homozygous mutant males. This phenotype is the same as observed in the testes of MIS ligand/α-inhibin double homozygous mutant males (Matzuk et al. 1995a).

Interestingly, unilateral stromal tumors were found in the testes of three MIS receptor homozygous mutants that were 5–7 months old. They also carried heterozygous mutations for cα-inhibin or MIS. The tumors differed slightly from those of inhibin mutants by being smaller, less hemorrhagic and containing nests of Leydig cells in addition to undifferentiated and granulosa cell-like foci (Fig. 4D). None of the six MIS receptor heterozygotes and α-inhibin wild-type mice in this series or any of the ~40 α-inhibin heterozygotes over 12 months of age in our earlier studies (Matzuk et al. 1995a) had stromal tumors. The mice mentioned above were examined on a C57Bl/6 × 129/SvEv mixed genetic background. The low penetrance of these stromal tumors in the MIS receptor homozygous mutant males may be caused by genetic background influences or stochastic mechanisms.

Taken together, these observations suggest that MIS signaling can influence the development of Sertoli/granulosa cell tumors initiated by the absence of inhibins and that inhibins can influence the development of Leydig cell neoplasia initiated by the absence of MIS signaling. The observation that the testicular tumors that developed in the MIS receptor/α-inhibin double homozygous...
mutant males were identical in phenotype to the testicular tumors that develop in the MIS ligand/α-inhibin double homozygous mutant males provides further evidence that MIS is the only ligand for the MIS receptor. In addition, these results suggest that the synergism between the MIS and inhibin signaling pathways occurs downstream of each receptor, not between ligands and receptors.

**Discussion**

*The MIS type II receptor gene*

Targeted mutagenesis of the MIS type II receptor gene leads to the development of males with internal pseudohermaphroditism, infertility, seminiferous tubule atrophy, and Leydig cell hyperplasia. Males that are otherwise normal have a uterus and oviducts, a phenotype that is identical to MIS ligand mutant male mice (Behringer et al. 1994). However, in contrast to MIS ligand mutant mice, MIS receptor mutants synthesize MIS ligand but possess target organs that are insensitive to its action. In addition, MIS ligand/MIS receptor double mutant males have the same mutant phenotype as either of the single mutants. Furthermore, MIS receptor/α-inhibin double mutant males develop dilated fluid-filled uteri and rapidly growing testicular tumors, a phenotype that is identical to male MIS ligand/α-inhibin double mutant mice [Matzuk et al. 1995a]. Female mice that lack MIS are normal and fertile as are the female MIS receptor mutant mice and the female MIS ligand/MIS receptor double mutant mice (Behringer et al. 1994). These genetic studies demonstrate that the MIS type II receptor gene functions in vivo to mediate MIS signaling.

In humans, the molecular basis of the PMDS syndrome is MIS receptor/α-inhibin double mutant male mice. Abdominal views of 4-month-old mutant male mice. (A) MIS receptor/α-inhibin double mutant; (B) MIS ligand/α-inhibin double mutant, (C) α-inhibin homozygous mutant. Note the larger testicular tumors (arrows) and expanded fluid-filled uteri in A and B. (D) Testes from 6-month-old male mice. MIS ligand-deficient testes were normal in size like the wild-type testis shown at the left. The α-inhibin-deficient testis on the right was enlarged and hemorrhagic because of an invasive Sertoli/granulosa tumor. The very large size of the two testes in the middle from MIS ligand/α-inhibin and MIS receptor/α-inhibin double-mutant mice were less hemorrhagic. In the testicular tumors of MIS receptor/α-inhibin double-mutant males, multifocal Leydig cell neoplasia and multifocal Sertoli/granulosa cell tumors were found that were identical to the tumors of ligand/α-inhibin double-mutant males. Bar, 1 cm.

Figure 5. Gross analysis of the reproductive tracts and testes of MIS receptor/α-inhibin double mutant male mice. Abdominal views of 4-month-old mutant male mice. (A) MIS receptor/α-inhibin double mutant; (B) MIS ligand/α-inhibin double mutant, (C) α-inhibin homozygous mutant. Note the larger testicular tumors (arrows) and expanded fluid-filled uteri in A and B. (D) Testes from 6-month-old male mice. MIS ligand-deficient testes were normal in size like the wild-type testis shown at the left. The α-inhibin-deficient testis on the right was enlarged and hemorrhagic because of an invasive Sertoli/granulosa tumor. The very large size of the two testes in the middle from MIS ligand/α-inhibin and MIS receptor/α-inhibin double-mutant mice were less hemorrhagic. In the testicular tumors of MIS receptor/α-inhibin double-mutant males, multifocal Leydig cell neoplasia and multifocal Sertoli/granulosa cell tumors were found that were identical to the tumors of ligand/α-inhibin double-mutant males. Bar, 1 cm.
Negative cases, MIS serum levels are low or undetectable, biopsied testicular tissue has no anti-Müllerian activity and mutations of the MIS gene have been detected (Knebelmann et al. 1991; Carré-Eusèbe et al. 1992; Imbeaud et al. 1994). In contrast AMH-positive patients have serum MIS levels at the upper limit of normal and testicular biopsies are able to induce the regression of fetal rat Müllerian ducts in culture. Testicular biopsies of PMDS dogs also exhibit anti-Müllerian activity (Meyers-Wallen et al. 1989). The insensitivity of MIS target organs to normal levels of the hormone suggests that a mutation of the MIS receptor could be involved and indeed, several have been detected by single strand conformation polymorphism (SSCP) screening in AMH-positive patients (Imbeaud et al. 1995, 1996). Interestingly, patients with MIS gene and MIS type II receptor gene mutations shared the same clinical phenotype. Thus, the mouse and human studies complement each other, and suggest that in mammals the MIS signaling pathway is simple.

The simplicity of the MIS signaling pathway relative to the complexity of other TGF-β family signaling pathways

In mammals, the MIS ligand mutant male phenotype is a phenocopy of the MIS type II receptor mutant male phenotype (Knebelmann et al. 1991; Behringer et al. 1994; Imbeaud et al. 1994, 1995, 1996). These findings are in contrast to those of other TGF-β family ligands and receptors whose genes have been mutated in mice. Activins are composed of βA and βB subunits (Vale et al. 1990). Activin-βA mutant mice are viable with defects in eyelid development and female reproduction (Vassali et al. 1994). Activin-βA mutant mice die within 24 hours after birth, lack whiskers and the lower, incisors and also have defects in the secondary palate, including cleft palate (Matzuk et al. 1995b). Activin βA/βB double mutant mice have also been generated that cannot produce activins or inhibins (Matzuk et al. 1995b). The mutant phenotype of these activin βA/βB double mutant mice is the addition of the activin-βA and activin-βB mutant phenotypes. In contrast, most activin type II receptor mutant mice are viable (Matzuk et al. 1995c). A minority of the activin receptor mutants have a variably expressed hypoplasia of the mandible that secondarily leads to an absence of incisors and cleft palate. These animals also have eyelid closure defects. The surviving male activin receptor mutants are fertile, but they are delayed in their ability to generate sufficient amounts of functional sperm. Female activin receptor mutants are infertile and have no estrous cycle. Significantly, few of the activin type II receptor mutant phenotypes overlapped with the mutant phenotypes of the activin ligand mutants, suggesting that in mammals, activin signaling pathways are complex.

Somewhat similar findings have been observed for BMP ligand and receptor mutant mice. BMP-2 mutant mice die during embryogenesis with a variable mutant phenotype. The predominant abnormality is a failure of the proamniotic canal to close. Subsequently, heart abnormalities also develop (Zhang and Bradley 1996). BMP-4 mutant mice die during embryogenesis with a highly variable phenotype, that is, some BMP-4 mutants can develop to the early somite stage whereas the most severely affected ones are arrested at the egg-cylinder stage and produce some extraembryonic mesoderm (Winnier et al. 1995). BMP-2/BMP-4 receptor mutant embryos do not form any mesodermal tissues and die by E8.5. In contrast with the variable mutant phenotypes of the BMP-2 and BMP-4 ligand mutants, the phenotype of the BMP-2/BMP-4 type I receptor mutant mice is consistent even when examined on multiple genetic backgrounds (Mishina et al. 1995). Although the most severe phenotype of the BMP-4 mutants approximate the BMP-2/BMP-4 type I receptor mutant phenotype, it is not a phenocopy.

Thus, in mammals at least the activin and the BMP-2/BMP-4 signaling pathways are complex. This may not be surprising because there are multiple type I and type II receptors for activins and BMP-2 and BMP-4 that have been defined by biochemical studies (Kingsley 1994). Furthermore, the contrasting findings between activin and BMP-2 and BMP-4 ligand mutant mice and mice mutant for their respective receptors may also be attributable in part to crosstalk between ligands and receptors. TGF-β family ligands are structurally related to each other (Kingsley 1994) and it has been shown that one type of receptor can interact with several classes of ligands (Attisano et al. 1993; Koening et al. 1994; Suzuki et al. 1994; ten Dijke et al. 1994, 1995). In addition, other studies indicate that one ligand can interact with several classes of receptors (Attisano et al. 1992, 1993; ten Dijke et al. 1994). As clearly shown here, unlike the complexity of other TGF-β family signaling pathways, the MIS signaling pathway is simple with MIS being the only ligand which interacts with the MIS type II receptor.

The role of MIS signaling in Müllerian duct regression and the testis

Although the MIS type II receptor is expressed in both testes and ovaries (Baarends et al. 1994, 1995a,b; di Clemente et al. 1994; Teixeira et al. 1996), our loss-of-function studies for MIS and the MIS receptor show that MIS signaling is not essential for either gonadal or germ cell development (Behringer et al. 1994). Whereas signaling through this receptor is essential for the regression of the Müllerian ducts during male sexual differentiation, these findings convincingly show that in the mouse, MIS signals are not essential for testicular morphogenesis and descent, and male and female gametogenesis. Thus, the perinatal loss of germ cells and subsequent loss of the ovary that was observed in female transgenic mice that overexpress hMIS may be caused by a crosstalk between MIS and other TGF-β family receptors. Alternatively, it is possible that the MIS type II receptor itself may interact with high levels of MIS to transduce an abnormal...
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signal to cause gonadal and/or germ cell abnormalities. The generation of transgenic mice that overexpress MIS in a MIS receptor mutant genetic background may provide clues to distinguish between these two possibilities.

A significant proportion of MIS receptor mutant males develop focal atrophy of the germinal epithelium associated with Leydig cell hyperplasia in the testis at older ages. In some cases, tubule atrophy was observed that was not associated with Leydig cell hyperplasia. Because MIS type II receptor expression has been shown only in Sertoli cells within the testis, the simplest interpretation of these results is that autocrine signaling by MIS would regulate Sertoli cell factors acting on neighboring Leydig and germ cells. MIS ligand deficient mice have Leydig cell hyperplasia and though not mentioned previously, also focal atrophy of the germinal epithelium (Behringer et al. 1994). Reexamination of nine MIS ligand mutant males older than 1.5 months detected this pathology in three of the animals. These results suggest a continuing role for MIS signals in the function of the testis during the period of reproduction.

Previously, we showed that the MIS and the inhibin signaling pathways can synergize to influence testicular tumorigenesis by generating mice that lacked both ligands (Matzuk et al. 1995a). It was not clear at what level MIS and inhibins were synergizing to influence tumorigenesis. The results presented here suggest that the synergism between the MIS and inhibin signaling pathways is not caused by interactions between inhibins and the MIS type II receptor. Therefore, the interaction between these two signaling pathways must be downstream of the receptors. Indeed, the interaction may be between different testicular cell types.

The MIS type I receptor and downstream targets of MIS signaling

TGFβ family receptor complexes are heteromers that contain type I and type II receptor molecules (Attisano et al. 1993; Franţen et al. 1993; Wrana et al. 1994; Franţen et al. 1995). Therefore, there should be a type I receptor that complexes with the MIS type II receptor to transduce signals that are essential for Müllerian duct regression and normal testicular function. Because the MIS type II receptor is only detected in MIS target organs, all tissue specificity for the action of MIS can be accounted for by the expression of the type II receptor. Therefore, although the MIS type I receptor must at least be expressed in the same cell types that express the MIS type II receptor, the MIS type I receptor may be more widely expressed and shared with other TGF-β family type II receptors.

Recently, a type I TGF-β family receptor has been isolated called ActR-1, Tsk7L, SKR1, ALK-2 or R1 that can bind both activin and TGF-β (Ebner et al. 1993; He et al. 1993; Matsuaki et al. 1993; ten Dijke et al. 1993). This type I receptor is expressed widely during embryogenesis and in adult tissues (He et al. 1993; ten Dijke et al. 1993). During rat embryogenesis, it is expressed in the mesenchymal cells adjacent to the Müllerian duct epithelium, suggesting that it might function as a MIS type I receptor.

Ultimately, it will be very interesting to determine the downstream targets of the MIS signaling pathway for Müllerian duct regression. Humans with PMDS who are MIS positive but have no apparent alterations in their MIS type II receptors may have mutations in genes encoding downstream effectors of MIS signaling. Indeed, whereas in humans, MIS maps to chromosome 19 (Cohen-Haguenauer et al. 1987) and the MIS type II receptor maps to chromosome 12 (Imbeaud et al. 1995; Visser et al. 1995), X-chromosome-linked transmission of PMDS has been reported in two families (Sloan and Walsh 1976; Naguib et al. 1989). Therefore, these two PMDS cases may be caused by a mutation in a third gene possibly functioning downstream of MIS. These human families and the MIS ligand and MIS type II receptor mutant mice will serve as valuable genetic resources to isolate the downstream targets of this differentiation pathway of male sexual development.

Materials and methods

Mutagenesis of the MIS type II receptor gene in mouse ES cells

A 129/SvEv mouse genomic library was screened with the extracellular-encoded region of the rabbit MIS type II receptor cDNA (di Clemente et al. 1994). Four positive λ-phage clones were isolated, and the genomic organization of the MIS type II receptor was characterized [Y. Mishina, R. Tizzard, B. Pathak, N. Copeland, N. Jenkins, R. Cate, and R. Behringer, in prep.]. A 3.9-kb HindIII–EcoRV 5′ fragment and a 4.2-kb XbaI–HindIII 3′ fragment were used to construct a replacement gene targeting vector [Fig. 1A]. A PGKneoβA resistance expression cassette was inserted in reverse orientation relative to the direction of the MIS receptor transcription between the two MIS receptor homologous regions (Soriano et al. 1991). An MC1tkpA herpes simplex virus thymidine kinase expression cassette was added onto the short arm of homology to enrich for homologous recombination by negative selection with 1-(2-deoxy-2-fluoro-β-D-arabinofuranosyl)-5-iodouracil (FIAU) [Mansour et al. 1988]. The targeting vector can be linearized at a unique NotI site outside of the homology. Twenty-five micrograms of linearized targeting vector was electroporated into 107 AB-1 ES cells that were subsequently cultured in the presence of G418 and FIAU on mitotically inactivated STO fibroblasts (McGahan and Bradley 1990, Soriano et al. 1991). Four hundred G418/FIAU-resistant ES clones were initially screened by SacI digestion and hybridized with a unique 3′ probe external to the region of vector homology. Correctly targeted clones were then expanded for further Southern blot analysis by EcoRI–EcoRV digestion and hybridization with a unique 5′ probe external to the vector homology. Two correctly targeted ES clones were identified.

Generation of chimeric mice and germline transmission of the MIS type II receptor mutant allele

Two of the MIS type II receptor mutant ES clones were microinjected into C57BL/6J blastocysts, and the resulting chimeric embryos were transferred to the uterine horns of day 2.5 pseudopregnant foster mothers (Bradley 1987). Chimeras were identified among the resulting progeny by their agouti fur [ES derived] and were subsequently bred with C57BL/6J mates. Two of the mutant ES clones [E8 and H8] were found to be capable of...
contributing to the germlines of chimeric mice. Tail DNA from the agouti pups that resulted from these matings was analyzed by Southern blot with either of the probes used to identify the MIS receptor heterozygotes. Chimeras were also bred with 129/SvEv females to establish the MIS receptor mutation on a 129/SvEv inbred genetic background.

**Generation of MIS ligand/MIS receptor double mutant mice and α-inhibin/MIS receptor double mutant mice**

To generate MIS ligand/MIS receptor double mutant mice, MIS ligand mutant mice were first bred with MIS receptor mutants to generate mice heterozygous for both mutations. The MIS ligand genotype was determined as described [Behringer et al. 1994]. These double heterozygous mice were interbred to generate mice homozygous mutant for both genes. Similarly, to generate α-inhibin/MIS receptor double mutant mice, α-inhibin mutant mice [Matzuk et al. 1992] were bred with MIS receptor mutants to generate mice heterozygous for both mutations. The α-inhibin genotype was determined as described [Matzuk et al. 1992]. These double heterozygous mice were interbred to generate mice homozygous mutant for both genes.

**Histological analysis**

Male reproductive tracts and testes were processed for histological analysis as described by Kaufman [1990]. Briefly, the tissues were fixed in 10% buffered formalin, dehydrated, and embedded in paraffin. Four-micrometer-thick sections were cut and stained with hematoxylin and eosin.

**MIS RT–PCR analysis**

Total RNA was isolated from the gonads of E13.5 embryos by the acid-guanidinium thiocyanate-phenol-chloroform method [Chomczynski and Sacchi 1987]. RNA was reverse transcribed with oligo(dT) 12-18 mer primer and M-MLV reverse transcriptase (Gibco BRL) at 25°C for 10 min and 42°C for 50 min. The resulting cDNA was amplified by PCR with the MIS specific primers 5'-GTCGAAACGCTATGAGTAC-3' and 5'-CCGCTGGTCCAGTATAAGC-3' and the HPRT specific primers 5'-CTCGTGACCTATTGACACT-3' and 5'-GCTCAGCCATAACCACAAA-3' and the reaction products were separated on 3% agarose gels and visualized by ethidium bromide staining.

**Measurement of serum MIS**

Serum MIS was measured by ELISA. Briefly, polystyrene plates Immulon-II (Dynatech, Guyancourt, France) were coated with an anti-recombinant human MIS polyclonal antibody raised in rabbit at 10 μg/ml. Serum was incubated at four different dilutions (1:4, 1:8, 1:16, and 1:32) in phosphate-buffered saline containing 1% BSA (PBS-1% BSA) for 1 h. The plates were successively exposed to an anti-recombinant human MIS polyclonal antibody raised in guinea pig at 5 μg/ml in PBS-1% BSA and to an alkaline phosphatase-conjugated anti-guinea pig IgG antibody [Rockland, Gilbertsville, PA] diluted 1:1000 in Tris-buffered saline containing 1% BSA. A color reaction was demonstrated with p-nitrophenyl phosphate [Sigma, St. Louis, MO] 1 mg/ml in diethanolamin/HCl 10 mM, MgCl₂, 1 mM at pH 9.8. Absorbance at 405 nm was read in a MR 700 spectrophotometer [Dynatech]. Data were processed by Biolinx software version 2.20 [Dynatech]. Calibration curves were constructed by use of increasing concentrations (0.44–27.5 ng/ml) of recombinant mouse MIS.

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