Xenopus embryos contain a somite-specific, MyoD-like protein that binds to a promoter site required for muscle actin expression

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We identify the "M region" of the muscle-specific Xenopus cardiac actin gene promoter from −282 to −348 as necessary for the embryonic expression of a cardiac actin–β-globin reporter gene injected into fertilized eggs. Four DNA-binding activities in embryo extracts, embryonic M-region factors 1–4 (EMF1–4), are described that interact specifically with this region. One of these, EMF1, is detected in extracts from microdissected somites, which differentiate into muscle, but not in extracts from the adjacent neuroectoderm, which differentiates into a variety of other cell types. Moreover, EMF1 is detected in embryo animal caps induced to form mesoderm, which includes muscle, and in which the cardiac actin gene is activated, but not in uninduced animal caps. EMF1 is also first detectable when cardiac actin transcripts begin to accumulate; therefore, both its temporal and spatial distributions during Xenopus development are consistent with a role in activating cardiac actin expression. Two lines of evidence suggest that EMF1 contains the myogenic factor Xenopus MyoD (XMyoD): (1) XMyoD synthesized in vitro can bind specifically to the same site as EMF1; and (2) antibodies raised against XMyoD bind to EMF1. DNA-binding studies indicate that EMF1 may be a complex between XMyoD and proteins found in muscle and other tissues. Our results suggest that the myogenic factor XMyoD, as a component of somite EMF1, regulates the activation of the cardiac actin gene in developing embryonic muscle by binding directly to a necessary region of the promoter.

[Key Words: MyoD; cardiac actin expression; myogenesis; Xenopus embryos; somites]

Received January 21, 1991; revised version accepted April 25, 1991.

In vertebrates, the cardiac actin gene is coexpressed with the skeletal actin gene in both the developing somites and the heart of the early embryo (Minty et al. 1982; Mohun et al. 1984; Sassoon et al. 1988). The focus of this study is cardiac actin expression in the somites. Muscle-specific expression requires only several hundred base pairs of the cardiac actin gene promoter (Minty and Kedes 1986; Mohun et al. 1986; Quitschke et al. 1989). One approach to defining the regulatory promoter sequences within this region and to identifying the transcription factors that interact with them has been to use cultured myogenic cell lines as a model for muscle differentiation. However, several investigators have noted anomalies in the patterns of gene expression shown by such established cell lines (e.g., Muscat and Kedes 1987; Vaidya et al. 1989; Braun et al. 1990), and it is uncertain how far these lines reflect the events of normal myogenesis.

An alternative approach is a direct analysis of gene activity in normal developing muscle during embryogenesis. In the frog, Xenopus laevis, expression of the cardiac actin gene is first detectable during gastrulation and is limited to that part of the newly formed mesoderm that gives rise to the somites (Mohun et al. 1984; Cascio and Gurdon 1986; Hopwood et al. 1989a). Its expression is maintained during the formation of the somites, which develop into the skeletal musculature. Cloned cardiac actin promoter–reporter gene constructs microinjected into fertilized eggs are expressed in a similarly restricted manner (Mohun et al. 1986; Wilson et al. 1986), which provides a direct assay to identify the promoter regions responsible for the embryonic activation of the cardiac actin gene.

Like all vertebrate muscle actin promoters (Minty and Kedes 1986; Mohun et al. 1986, 1989), the Xenopus cardiac actin promoter contains multiple copies of the 10-bp CArG box motif, CC(A/T)6GG. Mutational analysis revealed that the most proximal of the four CArG boxes, CArG box1, which is centered at −85 (Fig. 1A), is required for promoter activity in the frog embryo (Mohun et al. 1989). CArG box1 binds the ubiquitous transcription factor, serum response factor (SRF) (Taylor et al. 1989, Mohun et al. 1991), and at least three other Xenopus embryo proteins (Taylor 1991). Although the CArG
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box, as well as its flanking sequences, is sufficient for muscle-specific expression of the chicken skeletal actin gene in primary cell cultures [Walsh 1989], it is neither sufficient for high-level expression directed by the human cardiac actin promoter in myogenic cell lines [Minty and Kedes 1986, Sartorelli et al. 1990], nor sufficient for activation of the Xenopus cardiac actin gene in frog embryos [Mohun et al. 1986, 1989].

In previous studies we found that an internal deletion of the Xenopus cardiac actin promoter that left CArG box1 intact, but that removed the other three CArG boxes positioned between −104 and −224, had no inhibitory effect on the level or specificity of reporter gene expression in embryos. However, truncation of the promoter from −417 to −218 abolished transcription, indicating that sequences within this distal region were necessary for embryonic expression (Mohun et al. 1986, 1989).

In this paper we localize an essential distal sequence to a region from −282 to −348 in the cardiac actin gene promoter and show that it can bind the frog homolog of the mouse myogenic factor MyoD [Davis et al. 1987; Hopwood et al. 1989b, Harvey 1990; Scales et al. 1990]. We describe four sequence-specific DNA-binding activities in the normal developing embryo that interact with this promoter region. One of these is first detected during early neurulation and is detectable in extracts from somites but not from nonmuscle tissues. It also contains Xenopus MyoD (XMyoD) or a related protein. The XMyoD gene is expressed in the somitic mesoderm before the activation of the cardiac actin gene (Hopwood et al. 1989b; Harvey 1990; Scales et al. 1990), and our data suggest a simple model in which XMyoD regulates the activation of the cardiac actin gene in embryonic somites by binding directly to a necessary region of the promoter.

Results

A cardiac actin promoter sequence required for gene activation in embryos

Previously, we defined two regions of the cardiac actin

Figure 1. The M region is required for expression in embryos directed by the cardiac actin promoter. (A) Diagram of the Xenopus cardiac actin gene promoter (−417 to +1) showing the transcription initiation site (large arrow), TATA box (open rectangle), four CArG boxes (solid ovals), the M region (cross-hatched rectangle), and the series of deletions made in the promoter to identify essential distal regulatory sequences. (B) The effect of promoter deletions on transcription of a Xenopus cardiac actin–human β-globin fusion gene injected into fertilized Xenopus eggs. The expression of each deletion construct (test) was compared with that obtained from a similar fusion gene (ref) containing the complete cardiac actin promoter extending to −580 attached to a truncated human β-globin transcription unit. RNA from cultured embryo fragments was analyzed by RNase protection to detect transcripts (arrowhead) from the injected genes (actin–globin) and from the endogenous cardiac actin gene. (M) pBR HindIII molecular weight markers; (P) RNA probe; (t) tRNA control. Animal caps comprising prospective epidermis were cultured in isolation (−) or induced by conjugation with vegetal pieces to form axial structures, including embryonic muscle (+). Lanes 5 and 6 are from a longer exposure of the autoradiograph, which reveals a low level of residual probe. (C) The nucleotide sequence from −351 to −296 contains three E-box motifs [underlined] and corresponds to M3. One 20-nucleotide region shown separately as M1, is centered on a 14-nucleotide sequence, which is similar to the consensus-binding site proposed for MyoD.
promoter that are essential for expression of the gene in *Xenopus* embryos: a CArG box, from −80 to −90, and a more distal region, lying between −218 and −417 [Mohun et al. 1986, 1989]. To identify the necessary sequences within this distal region, we tested the effect of internal deletions on transcription directed by the cardiac actin promoter [Fig. 1A].

Chimeric *Xenopus* cardiac actin–human β-globin fusion genes were injected into fertilized *Xenopus* eggs, and their expression was monitored by an RNase protection assay during differentiation of embryonic muscle. To estimate the quantitative effect of each deletion, a reference fusion gene containing an unmodified cardiac actin gene promoter was coinjected with the test gene (see Materials and methods). The specificity of transcription from the injected genes was assayed by an embryologic manipulation that we have used previously. Animal caps dissected from blastula embryos can be induced by vegetal pieces to differentiate into axial structures, a major one of which is muscle [Sudarwati and Nieuwkoop 1971], and express both endogenous and microinjected cardiac actin genes [Gurdon et al. 1985; Mohun et al. 1986, 1989]. This manipulation is thought to mimic the events of mesoderm formation in normal development. In this study we compared RNA from these animal–vegetal conjugates with that from animal caps alone, which form atypical epidermis and do not express the cardiac actin gene.

Removal of the region from −104 to −224 placed the distal portion of the promoter (−225 to −417) directly adjacent to CArG box1 and did not reduce expression of the test fusion gene [Fig. 1B, lanes 3 and 4] [Mohun et al. 1989]. Similarly, increasing the extent of the internal deletion to encompass −104 to −282 left expression unaffected [lanes 7 and 8]. When the deletion extended to −348, however, expression of the fusion gene was virtually undetectable [lanes 9 and 10], indicating that a 3′ boundary for essential sequences within this portion of the promoter lies between −282 and −348. Furthermore, deletion of the region from −275 to −348 from the unmodified promoter was also sufficient to reduce dramatically expression of the fusion gene [lanes 5 and 6] and confirmed the importance of this region. We refer to the sequence from −282 to −348 as the “M-region” of the promoter.

**Temporal and spatial distribution of M-region-binding activities during embryogenesis**

We then used an electrophoretic mobility-shift assay [EMSA] to determine whether *Xenopus* embryos contain sequence-specific DNA-binding proteins that interact with the M region. An oligonucleotide probe (M3) comprising almost the entire M region [−351 to −296; Fig. 1C] was incubated with a whole-cell extract from stage-18 neurula embryos, a stage when embryonic muscle has begun to differentiate and the cardiac actin gene is expressed. Four of the protein–M3 DNA complexes detected were shown to be specific by competition with unlabeled M3 oligonucleotide, but not with the unrelated CArG box1 oligonucleotide [Fig. 2A]. We have called them EMF1–4 for embryonic M-region factors. EMF2–4 are readily detected, whereas EMF1 is a minor component that migrates just above EMF2 and is poorly resolved in this experiment [but see Fig. 2B, lanes 5–7].

Figure 2B shows these binding activities in a series of extracts from successive stages of early development. EMF1 is undetectable in egg extracts but detectable in embryo extracts from the early neurula stage onward. EMF3 and EMF4 are detectable at low levels in the egg, and their levels increase during early development. The level of EMF2 also increases somewhat, but other experiments indicate that any increase is comparatively small. In contrast, four proteins, including SRF, that bind specifically to CArG box1 are detected at an approximately constant level throughout this period [Mohun et al. 1989; Taylor 1991].

Having analyzed the temporal pattern of specific M3-binding activities, we investigated their spatial distribution. In neurula embryos, cardiac actin expression is restricted to the myotomes, which make up almost all of the somites in *Xenopus*. We used an extract made from somites dissected from late neurula in an EMSA with the M3 probe. EMF1–4 were detected, and each was competed by unlabeled M3 DNA (Fig. 2C). The sensitivity of the assay was such that extract from the somites of only two embryos was sufficient to generate a signal. EMF1 was relatively prominent in the somite extract compared with whole embryo extracts, suggesting that it is enriched in embryonic muscle cells.

Two lines of evidence support this view. First, EMF1 is undetectable in extracts of dissected neurectoderm, a nonmuscle tissue adjacent to the somites that differentiates to form a range of nonmuscle cell types, including nervous tissue and epidermis [Fig. 2C, cf. lanes 1 and 2]. This contrasts with the distribution of the other bands, which are of similar intensity with each extract. The difference between the extracts is most striking when the detection of EMF1 is enhanced by omission of Mg2+ from the assay [Fig. 2C, lanes 3 and 4]. Under these conditions, EMF2, which migrates closely to EMF1, binds poorly. Second, EMF1 is undetectable in extracts of belly pieces from stage-34 tadpoles [data not shown]. These pieces contain a different range of nonmuscle cell types than those found in the neurectoderm. In summary, both the temporal and spatial distributions of EMF1 during *Xenopus* embryogenesis are consistent with a role for it in activating expression of the cardiac actin gene.

**Xenopus MyoD can bind to the M region**

Examination of the M region sequence revealed three copies of the E-box motif, CANNTG [see Fig. 1C], which is common to the binding sites of several helix–loop–helix (HLH) DNA-binding proteins [Murre et al. 1989a] and lies at the center of the mouse MyoD-binding site in the muscle creatine kinase (MCK) enhancer [Lassar et al. 1989]. We therefore asked whether *Xenopus* MyoD could bind to the M region. XMMyoD protein prepared by translation in vitro of synthetic XMMyoD RNA was used in an...
EMSA with the M3 probe. XMyoD does bind to M3 (Fig. 3, lane 5). The XMyoD–M3 interaction is specific for both the protein and the DNA probe. First, no specific complex is produced with either mock-translated reticulocyte lysate or with in-vitro-translated *Xenopus* SRF [lanes 1 and 2], which binds to cardiac actin CArG box1. Second, formation of the M3–XMyoD complex is competed by unlabeled M3 oligonucleotide, but not by CArG box1 [lanes 6, 7, and 9]. The amount of complex formed on the M3 probe is similar to that formed in a binding assay with the same amount of XMyoD and an equivalent amount of probe containing the two MyoD-binding sites of the mouse MCK enhancer [Fig. 3, lanes 10–12]. This indicates that the affinity of XMyoD is similar for each sequence.

The binding of XMyoD to M3 requires an intact first helix in its HLH domain, which suggests that XMyoD can bind as a homodimer to M3. The mutant XMyoD114P [Hopwood and Gurdon 1990] has a single-amino-acid substitution that disrupts the putative HLH domain, which in other systems is required for MyoD to dimerize with itself and other proteins and for subsequent DNA binding [Murre et al. 1989b; Davis et al. 1990]. No specific complex was detectable using the same amount of XMyoD114P protein as was used in the assays with wild-type XMyoD [Fig. 3, lane 4].

The other member of the MyoD family characterized thus far in *Xenopus* is a probable homolog of the human gene Myf-5 [Hopwood et al. 1991]. XMyoD and XMyf5 proteins are closely related; therefore, we asked whether XMyf5 could also bind to the E boxes in the M region of the cardiac actin promoter. In contrast to the results obtained with XMyoD, no specific DNA–protein complex was detected when in vitro-translated XMyf5 was used in an EMSA with M3 DNA probe [Fig. 3, lane 3]. XMyf5 can, however, participate in binding to M3 in combination with other proteins (see below).

**XMyoD and EMF1 both bind to a 20-bp sequence within the M region**

Of the three E-box motifs within the M3 sequence, only one shows extensive similarity to the 14-bp consensus originally proposed for the MyoD-binding site [Buskin and Haushka 1989] and to the shorter consensus for MyoD–E2A heterodimers recently defined by the selection of binding sites from random oligonucleotides [Blackwell and Weintraub 1990]. A 20-bp oligonucleotide [M1; see Fig. 1C] that encompasses this sequence binds XMyoD specifically [Fig. 4A, lanes 1–5], and is an effective competitor of XMyoD binding to M3 [Fig. 3, lane 8]. The affinity of XMyoD protein is similar for both M1 and M3 [cf. Fig. 3, lane 5 with Fig. 4A, lane 3], indicating that the M1 sequence is the major component of the XMyoD-binding activity of M3. However, dimethylsulfate interference assays using the M3 probe and XMyoD–GST (glutathione-S-transferase) fusion protein show that in addition to binding at the M1 site, XMyoD can also...
might be a component of the somite EMF1 complex. We tested the possibility that somite EMF1 contains an XMyoD-related protein by using an antiserum raised against XMyoD-GST fusion protein in an EMSA with the somite extract. Two approaches were used with both the M3 and M1 probes. In the first, a “supershift” assay, the antibodies were added after the formation of the EMF1–M region DNA complex; and in the second, a depletion assay, the antibodies were added to the somite extract before the M region DNA. The EMF1 complex was completely supershifted into a slower migrating band by the anti-XMyoD–GST serum [lanes 1–4]. Similarly, the EMF1 complex was completely removed in the depletion assay by both the anti-XMyoD–GST serum and anti-XMyoD antibodies affinity-purified from this serum [lanes 5–10]. In each approach the specificity of the antibody action was demonstrated by no effect of both pre-immune serum [lanes 1, 3, 5, and 8] and anti-GST antibodies [not shown].

In whole-stage 18 embryo extracts, EMF1 was only just detectable (see Figs. 2B and 4B), but here too it was supershifted by the anti-XMyoD antibodies [not shown]. No other complex, including EMF2–4 and any detected with the neuroectoderm extract, was affected by inclusion of these antibodies in the binding reaction, indicating that they bound specifically to a component of the EMF1 complex. In addition, the anti-XMyoD antibodies did not cross-react with an XMyf5-containing complex (see below). We conclude that EMF1 contains XMyoD or an antigenically related protein.

**EMF1 contains an XMyoD-related protein**

We tested the possibility that somite EMF1 contains XMyoD by using an antiserum raised against XMyoD–GST fusion protein in an EMSA with the somite extract. Two approaches were used with both the M3 and M1 probes [Fig. 5]. In the first, a “supershift” assay, the antibodies were added after the formation of the EMF1–M region DNA complex; and in the second, a depletion assay, the antibodies were added to the somite extract before the M region DNA. The EMF1 complex was completely supershifted into a slower migrating band by the anti-XMyoD–GST serum [lanes 1–4]. Similarly, the interaction with the adjacent, upstream E box [data not shown].

Using the same 20-bp M1 probe, we detected a specific M1-binding activity [arrowhead] in embryo extracts with the temporal and spatial characteristics of the M3-binding activity EMF1 [Fig. 4B]. The M1-binding activity was detectable in the staged extracts from early neurula onward [lanes 1–7] and was present in the extracts from somites, but not those from neuroectoderm [lanes 8 and 9]. It was specific, as shown by competition with unlabeled M1 oligonucleotide [lanes 10–12]. In addition, M1 oligonucleotide competed formation of the somite EMF1–M3 complex [not shown]. These results identify a 20-bp-binding site of EMF1 and suggest that XMyoD might be a component of the somite EMF1 complex.

**Mixing XMyoD with neuroectoderm extract produces a binding activity that comigrates with EMF1**

The complexes of in vitro-translated XMyoD and the M region probes migrate substantially faster than the somite complex EMF1. We reasoned that if EMF1 contains XMyoD, it does so in combination with other proteins. Because the myogenic proteins can form heterodimers–hetero-oligomers with other HLH proteins in vitro (Murre et al. 1989b, Davis et al. 1990), it is possible that EMF1 is XMyoD complexed with another member of the HLH family. To investigate these other factors we tested whether nonmuscle extracts contain proteins that can reconstitute an EMF1-like-binding activity when mixed with XMyoD.

The addition of in vitro-translated XMyoD to neuroectoderm extract resulted in the appearance of an M region-binding activity that was undetectable with the un-supplemented neuroectoderm extract [Fig. 6A, lanes 1–4]. No such complex was formed with neuroectoderm extract mixed with mock-translated reticulocyte lysate [lane 8], and the complex was specific, as demonstrated by competition with unlabeled binding site oligonucleotide [not shown]. The new complex was also formed with a concentration of XMyoD below that required to detect binding of XMyoD alone [lanes 5–8], indicating that it had a binding affinity higher than XMyoD alone. The mobility of the complex was less than that of XMyoD alone [Fig. 6A, lanes 2 and 3] but, strikingly, was indistinguishable from that of somite EMF1 [Fig. 6B, lanes 1–4]. Similarly, mixing XMyoD with somite extract produced more of the same mobility complex [not shown]. These results suggest that EMF1 might be a complex between XMyoD and proteins that are not muscle specific. Association with these proteins requires an intact HLH domain, because XMyoD114P, which has a point mutation that disrupts the HLH motif, does not form the new complex when it is mixed with neuroectoderm extract [Fig. 6B, lanes 5 and 6].

Similar results were also obtained in mixing experiments using XMyf5 protein [Fig. 7] despite the lack of interacting with the adjacent, upstream E box (data not shown).

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Similar results were also obtained in mixing experiments using XMyf5 protein (Fig. 7) despite the lack of
detectable binding of XMyf5 alone to the M region. Thus, the complex between the neuroectoderm extract and XMyf5, as with XMyoD, has a higher binding affinity than the myogenic factor alone. Similarly, it also requires an intact HLH domain for its formation: No complex is detectable with XMyf5-102P, which has a point mutation in the HLH motif [Hopwood et al. 1991]. However, the XMyf5–neuroectoderm complex migrates more rapidly than both somite EMF1 and the EMF1-like activity formed by mixing XMyoD and neuroectoderm extract [Fig. 6B, lanes 1–4; Fig. 7, lanes 4 and 6]. This is consistent with predicted relative molecular masses of 32 kD for XMyoD and 28 kD for XMyf5 [Hopwood et al. 1989b, 1991].

We were then able to use the binding of XMyf5 to assess in the EMSA the specificity of the antibodies for XMyoD in comparison with XMyf5, the closest known relative of XMyoD in *Xenopus*. Incubation with anti-XMyoD–GST serum or the affinity-purified anti-XMyoD antibodies, but not preimmune serum or anti-GST serum (not shown), completely removed the band corresponding to the XMyoD-containing complex formed by mixing XMyoD with stage-18 embryo extract [Fig. 8, lanes 1–3]. In contrast, the band corresponding to the XMyf5-containing complex was unaffected by these anti-XMyoD antibodies [Fig. 8, lanes 4–6]. This demonstrates that under the conditions used in these assays, the antibodies do not cross-react with a complex containing XMyf5. The antibodies do, however, completely remove or supershift somite EMF1 [Fig. 5], which indicates that the complex formed between XMyf5 and embryo extract is not a detectable component of EMF1 (see Discussion). We conclude that somite EMF1 is a complex between XMyoD, or a similar protein, and another protein present in many, if not all, embryo tissues.

**The EMF1-binding activity is inducible in animal caps**

Animal caps can be diverted experimentally from their normal ectodermal pathway of differentiation to form mesodermal derivatives, including muscle. Extracts made from animal caps of stage-8 blastulae, which had been induced to form mesoderm by incubation in XTC-conditioned medium, were compared with extracts from uninduced animal caps in the EMSA with the M3 probe. An M3-binding activity, EMF1, was detectable in the extract from induced animal caps but was undetectable in the extract from uninduced animal caps [Fig. 9, lanes 1 and 2]. This binding activity was specific, as shown by competition with unlabeled M3 oligonucleotide [lane 5], and was identified as EMF1 as follows: It was removed by incubation of the extract with the anti-XMyoD antiserum, but not preimmune serum, before the addition of the M3 probe [lanes 3 and 4], and it comigrated in the EMSA with the previously defined EMF1 [not shown]. Similar results were obtained with extracts from animal caps induced by making conjugates with vegetal pieces of blastulae.
In this article we have shown that the M region from -282 to -348 of the cardiac actin promoter is required for expression of this gene in embryos. Synthetic XMyoD can bind specifically to this region, probably as a homodimer, with an affinity similar to that of the mouse MCK enhancer. We have extended these findings by analyzing Xenopus embryo factors that interact with this MyoD-binding site. Early frog embryos can be dissected into discrete tissues, and we have exploited this to describe four specific M-region-binding activities that are present in somites. Any of these could be required for cardiac actin expression. However, one of them, EMF1, is first detectable when cardiac actin transcripts begin to accumulate and appears to be restricted to embryonic muscle. This suggests a simple model in which EMF1 binding to the M region of the cardiac actin promoter results in tissue-specific activation of the gene.

Our results indicate that one component of EMF1 is XMyoD: [1] EMF1 is detectable in embryonic muscle but not in a variety of nonmuscle tissues; [2] EMF1, as with XMyoD [Hopwood et al. 1989b], is detectable in animal caps induced to form muscle but not in uninduced animal caps; [3] EMF1 is not detectable in UV-treated embryos [not shown], which contain no muscle [Scharf and Gerhart 1980] or XMyoD RNA [N.D. Hopwood and K. Kao, unpubl.]; [4] EMF1 is first readily detectable in early neurulae, and long exposures of autoradiographs indicate a low level in gastrulae; [5] EMF1 binds to a 20-bp promoter sequence containing a 14-bp MyoD-binding site consensus; and [6] anti-XMyoD antibodies bind to EMF1. Our results also suggest that EMF1 is an HLH domain-dependent complex, a dimer or higher order multimer, between XMyoD and other proteins found in nonmuscle tissues, as well as in somites. This protein could be a product of the Xenopus E2A gene [Murre et al. 1989a; Kamps et al. 1990]. Consistent with this idea, the complex between XMyoD and neurectoderm extract shares several properties with the MyoD–E12 complex [Murre et al. 1989b; Davis et al. 1990]: Both migrate more slowly and have a higher binding affinity than MyoD alone, and both can form post-translationally.

Experiments with cultured cells in which MyoD induced muscle differentiation in nonmuscle cell lines suggest that it plays a critical role in establishing muscle differentiation [Davis et al. 1987; Weintraub et al. 1989]. MyoD also appears to participate in the activation of several genes expressed during the terminal differentiation of muscle cells. Thus, the mouse MCK enhancer binds both synthetic MyoD and a complex from a muscle cell line that interacts with anti-MyoD antibodies [Lassar et al. 1989]. Moreover, this binding site is required for full enhancer activity [Buskin and Hauschka 1989; Lassar et al. 1989] and is sufficient for transcriptional activation by MyoD after transfection into C3H/10T1/2 fibroblasts [Weintraub et al. 1990]. Synthetic MyoD has also been shown to bind functionally to important sites in the acetylcholine receptor α-subunit gene promoter, the myosin light-chain 1/3 enhancer [Braun et al. 1990; Piette et al. 1990] and, most recently, the human cardiac actin promoter [Sartorelli et al. 1990]. In this last case, mutation of the MyoD-binding site, which lies between the TATA box and the most proximal CArG box, greatly reduces expression from the promoter in both muscle cell lines and nonmuscle cell lines supplied with MyoD. The site may therefore play an analogous role to the M region of the Xenopus gene, despite the difference in its location and the presence of only a single binding motif, which contrasts with the multiple sites in the Xenopus gene and the other muscle gene promoter–enhancers that bind MyoD [Lassar et al. 1989; Piette et al. 1990; Rosenthal et al. 1990].

Although much progress has resulted from studies of cultured cells, it is important to assess the role of MyoD during embryogenesis, when developmental fate decisions are made and muscle is differentiating normally. In the Xenopus embryo, XMyoD transcripts accumulate, above a low maternal level, in the presumptive somitic mesoderm of the gastrula ~2 hr before cardiac actin transcripts first appear [Hopwood et al. 1989b]. This pattern of expression is consistent with a function for XMyoD in the early events of myogenesis in the frog embryo, including the activation of the cardiac actin gene. This latter role is suggested by the finding that in the Xenopus embryo ectopic expression of XMyoD, but not the HLH domain mutant XMyoD114P, activates the cardiac actin
Figure 6. Mixing XMyoD with neuroectoderm extract produces an M region-binding complex that comigrates with EMF1. (A) EMSA with M3 probe and protein in each lane as follows: (1) Mock-translated reticulocyte lysate; (2) XMyoD; (3) XMyoD plus neuroectoderm extract; (4) neuroectoderm; (5) mock-translated reticulocyte lysate; (6) XMyoD; (7) XMyoD plus neuroectoderm; (8) mock-translated reticulocyte plus neuroectoderm. Lanes 5–8 have fivefold less XMyoD or control lysate. (Lower arrowhead) The XMyoD complex; (upper arrowhead) the complex obtained by mixing XMyoD with neuroectoderm extract. (B) EMSA with M1 probe and protein in each lane as follows: (1) XMyoD plus neuroectoderm extract; (2) somite extract; (3) XMyf5 plus neuroectoderm; (4) mock-translated reticulocyte lysate plus neuroectoderm; (5) XMyoD plus neuroectoderm; (6) XMyoD114P plus neuroectoderm. Complexes obtained by mixing neuroectoderm extract with XMyoD [upper arrowhead] and XMyf5 [lower arrowhead].

gene in cells that would not normally express it (Hopwood and Gurdon 1990). Somite-specific activation of the cardiac actin gene could therefore be a direct consequence of localized XMyoD expression in early somitic mesoderm. Such an effect might be direct through binding to the cardiac actin promoter, or indirect through the activation of other factors that regulate muscle differentiation. A major result of the work presented here is that normal embryo somites contain a MyoD-like factor that binds to the cardiac actin promoter in vitro. Moreover, in animal caps induced to form mesoderm in which transcription from the cardiac actin is strongly activated, the XMyoD-containing EMF1-binding activity is also induced. These results suggest a direct mechanism and a model in which XMyoD is the muscle-specific component of a DNA-binding complex that interacts with a site necessary for activity of the cardiac actin promoter.

The regulation of cardiac actin expression in *Xenopus* development is, however, likely to be more complex in several respects than this simple model suggests. First, other myogenic factors may have a role in activating cardiac actin expression. Ectopic expression of XMyf5, as with XMyoD, can activate the cardiac actin gene in *Xenopus* embryos (Hopwood et al. 1991); and we show here that XMyf5 can, in combination with other embryo proteins, bind to the M region. XMyf5 is probably not detected as a component of EMF1 as it is found mainly in posterior mesoderm at the stage when the somites were dissected (Hopwood et al. 1991), which would not have been included in our dissections. Second, although promoter elements other than the M region, including CArG box1, may not establish muscle-specific transcription, they may be required to sustain the high levels of transcription characteristic of differentiating muscle. Finally, in vertebrates the cardiac actin gene is coexpressed with its skeletal counterpart in both the developing heart and the axial musculature of the embryo (Minty et al. 1982; Mohun et al. 1984; Sassoon et al. 1988). Because MyoD is not detectable in either embryonic or adult heart (Davis et al. 1987; Hopwood et al. 1989b; Sassoon et al. 1989), a second, MyoD-independent mechanism must be responsible for activating the cardiac actin gene in cardiac muscle.

Materials and methods

Promoter analysis

Internal promoter deletions were introduced into the *Xenopus*
MyoD and embryonic muscle actin expression

Figure 7. Mixing XMyf5 with neuroectoderm extract produces an M-region-binding complex. EMSA is shown with M3 probe and protein in each lane as follows: (1) Mock-translated reticulocyte lysate; (2) XMyf5; (3) neuroectoderm extract. Lanes 4–7 are neuroectoderm extract plus (4) XMyf5; (5) XMyf5-102P; (6) XMyoD; and (7) mock-translated reticulocyte lysate. The binding activity produced by mixing XMyf5 with neuroectoderm extract is specific, as it is competed by unlabeled M3 oligonucleotide (not shown), and is dependent on XMyf5, as it is not formed by mixing mock-translated reticulocyte lysate with neuroectoderm extract. The complexes obtained by mixing neuroectoderm extract with XMyoD (upper arrowhead) and XMyf5 (lower arrowhead) are indicated.

Cardiac actin–human β-globin gene by recombining fragments from KpnI-linker scan mutations described previously (Mohun et al. 1989). Excision of the fusion gene as a Scal–HindIII fragment followed by recloning resulted in truncation of the deleted promoters from −580 to −417. Linearized test and reference plasmids (Mohun et al. 1989), 175 pg in 7 nl, were injected into each blastomere of two-cell embryos in a ratio of 5:1. At the blastula stage, animal pole explants were dissected and cultured either as pairs or as conjugates with vegetal portions of un.injected blastulae (Gurdon et al. 1985). Explants were frozen when sibling, un injected embryos had completed neurulation. RNA was extracted and analyzed as described previously (Mohun et al. 1989). In addition to protection by authentic transcripts from the injected genes, a number of bands are obtained that result from incomplete removal of injected DNA from the samples and indicate the presence of material in the assays of uninduced animal caps. The cardiac actin probe also detects transcripts from the endogenous gene: Two bands are revealed that result from the use of two polyadenylation signals (Mohun et al. 1986).

Probes

Oligonucleotidenucleotides for M3 and M1 (upper strands shown in Fig. 1C) were synthesized with KpnI-compatible ends and cloned into the KpnI site of the pUC18 polylinker. Probes were prepared by excising the insert with EcoRI and BamHI, and end-labeling with Klenow fragment and [α-32P]dATP. The CArG box I probe was as described (Mohun et al. 1991), and the MCK probe was made from the plasmid pUC-E (Buskin and Hauschka 1989) by exciting and end-labeling the 110-bp Aval–Neol fragment containing the two defined MyoD-binding sites of the MCK enhancer.

Embryo extracts

Embryos were staged according to Nieuwkoop and Faber (1967). The staged extracts were made from unfertilized eggs, and from morula (stage 6.5), blastula (stage 8–9), gastrula (stage 11–12), early neurula (stage 14–15), late neurula (stage 18), and tailbud (stage 24) embryos. The somite dissections, from stage 18–20 neurulae, comprised the anterior segmented region plus a similar amount of immediately posterior unsegmented somitic mesoderm. The corresponding neuroectoderm dissection comprised the adjacent neural tissue, including neural crest, together with the surrounding epidermis. The stage-34 belly piece was that remaining after the removal of the head, the axial and tail region, and the anterior ventral region containing the heart. Animal caps dissected from stage-8 blastulae were either cultured untreated or induced by culturing for 4 hr in a one-third dilution of heat-treated conditioned medium from XTC cells. Extracts were made when sibling embryos had completed neurulation. Whole-cell extracts were made on ice by homogenization of embryos or embryo pieces: 30 whole embryos in 300 μl, 20 somite/neuroectoderm dissections in 50 μl, 30 belly dissections in 55 μl, and 30 induced or uninduced animal caps in 45 μl of homogenization buffer (50 mM Tris-HCl [pH 7.9], 25% glyc-
translated XMyoD was detected more clearly without MgCl₂.
Protein [5 μl of embryo extract or ~60 fmoles of specific protein
{unless stated otherwise} made by in vitro translation] was
added last, the mix was incubated for 20 min at room temper-
ature, and the complexes were analyzed on 5% acryl-
amide/0.56× TBE gels. In the comparison of binding between
M1, M3, and MCK probes, the same amount of protein from the
same in vitro synthesis was used. The assays with in vitro-
translated XMyoD contained a 15-fold molar excess of protein
over DNA, compared with the 100~1000-fold molar excesses
used in many previous studies of other putative MyoD-binding
sites [Lassar et al 1989, Braun et al. 1990, Piette et al. 1990;
Rosenthal et al. 1990, Weintraub et al. 1990]. In assays with
mixed samples, embryo extracts were incubated with products
of in vitro translation for 5 min at room temperature before the
addition of probe. To make double-stranded competitor binding
sites, upper and lower strand gel-purified oligonucleotides were
annealed by heating to 65°C in 10 mM Tris-HCl (pH 8), 5 mM
MgCl₂, followed by slow cooling. Alternatively, competitor
DNAs were isolated by restriction enzyme digestion of the ap-
propriate plasmid and subsequent electrophoresis and elution of
the desired band. The assays in conjunction with antibodies
were as follows. A 5% dilution [2 μl] of unpurified antisemur or
affinity-purified antibodies was added to the protein mix and
incubated for 15 min on ice before the addition of DNA. The
incubation was continued for 15 min at room temperature be-
fore analysis by 5% acrylamide gel electrophoresis. For the
supershift assay the antisemur was added to the DNA–protein
complexes formed in the standard way and the incubation con-
tinued for 15 min on ice before analysis on a 3.5% gel.

Antibody production and purification

A plasmid for expression in Escherichia coli of XMyoD fused to
GST was made by cloning an AluI fragment of XMyoD2-24
{nucleotides 143~723} [Hopwood et al. 1989b] into the SmaI site
of pGEX-3X [Smith and Johnson 1988]. Protein for immuniza-
tion of rabbits was purified essentially as described [Smith
and Johnson 1988]. Anti-GST antibodies were removed by incuba-
tion with CNBr-activated Sepharose 4B coupled to GST, and
anti-XMyoD antibodies were purified by affinity chromatog-
raphy on CNBr-Sepharose 4B coupled to the XMyoD fusion
protein.

Acknowledgments

We thank Jean Buskin and Steve Hauschka for the pUC-E plas-
mid, Janet Heasman for XCT-conditioned medium and Kazuto
Kato and Ken Kao for advice on its use, Nick Torpey for anti-
GST antisemur, and Colin Sharpe and Jeremy Rashbass for com-
ments on the manuscript. We are grateful for the financial sup-
port of the Cancer Research Campaign, the Royal Society, and
the Medical Research Council.

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*Genes Dev.* 1991, 5: Access the most recent version at doi:10.1101/gad.5.7.1149

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