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Global transcriptional regulation of the locus encoding the skeletal muscle determination genes *Mrf4* and *Myf5*

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The linked *Mrf4* and *Myf5* genes encode two transcription factors essential for the determination and differentiation of skeletal muscle in the embryo. The locus is controlled by a multitude of interdigitated enhancers that activate gene expression at different times and in precisely defined progenitor cell populations. Manipulation of the enhancer–promoter composition of the locus reveals a novel mechanism for the regulation of such a gene cluster. Enhancers, promoters, and a new class of elements we call transcription balancing sequences, which can act as cryptic promoters, exist in a series of equilibria to ensure that enhancers and promoters together produce the highly dynamic and exquisitely specific expression patterns of the two genes. The proposed model depends upon nonproductive interactions between enhancers and both minimal and cryptic promoters, and is distinct from those developed for the β -*globin* and *Hox* clusters. Moreover, it provides an explanation for the unexpected phenotypes of the three *Mrf4* knockout alleles.

[*Keywords*: Transcriptional regulation; cryptic promoter; *Mrf4*; *Myf5*; myogenesis]

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Eukaryotes employ diverse mechanisms to control gene expression that can be categorized into transcriptional, post-transcriptional, and post-translational regulation. Of these, the rate of transcriptional initiation is probably the most important factor controlling the overall expression pattern of a gene. Basal promoters can initiate transcription on their own, but since most of the transcription factor-binding sites within them interact with ubiquitously expressed factors, transcription is not tissue-specific, and is usually driven at low levels. To increase transcription rates, and to confer specificity, enhancers interact with basal promoters by means that are not fully understood; the specificity is determined by the combination of binding sites in their sequences. In contrast, throughout mammalian genomes many cryptic promoters have been identified, sequences only able to initiate transcription following an alteration of their genomic environment, either experimentally or as the result of a translocation. These cryptic promoters must be prevented from interacting productively with the enhancers so that they remain transcriptionally silent. The expression pattern of any particular gene is established by the promoter/enhancer composition of the locus, the availability of cognate transcription factors, and the ki-

netics and mechanics affecting their interactions. The maintenance of the correct specificity of these interactions is crucial for orderly development and differentiation, and is a particular problem in loci containing multiple genes related by function and/or expression pattern. Of significant interest in this context is the regulation of genes encoding transcription factors that play key roles in a cascade controlling cell fate decisions.

During mouse embryogenesis, skeletal muscle arises from three different locations: segmented somitic paraxial mesoderm, unsegmented paraxial head mesoderm, and prechordal mesoderm. Trunk and limb muscles originate from the somitic epithelial dermomyotome (for review, see Kalchauer and Ben-Yair 2005). Progenitor cells delaminate from the four lips of the dermomyotome and move into the myotome, in which they develop in situ or subsequently travel to their final destination. At limb levels, cells from the ventrolateral lip (VLL) of the dermomyotome migrate into the developing limb buds, to form the appendicular musculature, while at more rostral positions VLL cells delaminate and follow a long-range migration along the hypoglossal chord to give rise to pharyngeal and tongue musculature. Facial musculature is derived from unsegmented head mesoderm and prechordal mesoderm cells, which either move through the branchial arches or directly to their final destination (for review, see Noden and Francis-West 2006).

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In all of these locations the determination, migration, and terminal differentiation of skeletal muscle cells are controlled by a network in which the four myogenic regulatory factors (MRFs)—*Myf5*, *Mrf4*, *MyoD*, and *myogenin*—play key roles. *Myf5* and *MyoD* can act as determination genes, while *myogenin* is essential for the terminal differentiation of committed myoblasts (for review, see Berkes and Tapscott 2005). *Mrf4* has long been considered a differentiation gene, but it is expressed in undifferentiated cells and can act as a determination gene (Kassar-Duchossoy et al. 2004)

We have focused on the transcriptional regulation of *Mrf4* and *Myf5*, which are separated by 8.7 kb in the mouse and are linked in all vertebrates. *Myf5* is the first MRF gene expressed in the mouse; it is activated before 8.5 d post-coitum (dpc) in the dorsomedial lip of the somite and then extends into the newly formed myotome; by 9.5 dpc, expression begins in the ventral dermomyotome and branchial arches. As limbs develop, *Myf5* is up-regulated in the central premyotome but not in migrating progenitors (Ott et al. 1991; Summerbell et al. 2000). During myoblast fusion expression is maintained in all skeletal muscle masses but it is down-regulated from late gestation, and only quiescent satellite cells and muscle spindles retain expression in the adult (Zammit et al. 2004). *Mrf4* expression closely follows that of *Myf5* but is triphasic. It begins in undifferentiated cells of the ventral somite coincident with the onset of *Myf5* expression in the dorsomedial lip (Carvajal et al. 2001; Summerbell et al. 2002). It is then activated in the central myotome of rostral somites at ~9.0 dpc, and by 10.0 dpc includes all somites but is absent from the limbs and branchial arch derivatives. This phase is down-regulated in a rostrocaudal direction in all somites from 12.5 dpc, followed by a third phase in which expression is up-regulated and maintained in all skeletal muscles in the adult.

The regulation of linked and functionally related genes has been studied in a number of systems. In the β -globin locus temporal switching from one gene to another in the same cell type is regulated by proximal sequences, insulators restrict interactions, while overall levels of expression are controlled by the distal locus control region (LCR), involved in chromatin architecture (for review, see de Laat and Grosveld 2003). In vertebrate *Hox* clusters global control regions regulate the expression of multiple genes, and locally acting elements shared between adjacent genes affect expression in particular cell types and tissues (Spitz et al. 2003). In *Drosophila* homeotic clusters, the precise structure of core promoter elements controls enhancer–promoter selectivity while a novel class of element (promoter tethers) facilitates interactions of enhancers with core promoters (Calhoun and Levine 2003).

Transcriptional regulation of *Myf5* has been studied extensively in transgenic mice (Patapoutian et al. 1993; Zweigerdt et al. 1997; Hadchouel et al. 2000, 2003; Summerbell et al. 2000; Carvajal et al. 2001; Teboul et al. 2002; Buchberger et al. 2003; Zammit et al. 2004). A similar approach has been used in the analysis of the transcriptional regulation of *Mrf4* (Patapoutian et al.

1993; Pin et al. 1997; Carvajal et al. 2001; Pin and Koniczny 2002). We have shown previously that a 250-kb BAC clone carrying 195 kb of sequence upstream of the *Myf5* translational start site (BAC195) is able to fully recapitulate the expression patterns of both *Mrf4* and *Myf5* during embryonic development (Carvajal et al. 2001) and in the adult (Zammit et al. 2004). A complex picture has emerged (Fig. 1A) in which a multitude of enhancers regulate the activation of *Myf5* at different times and anatomical locations during development, suggesting that the inductive signals for myogenesis are diverse, and that *Myf5* may function to canalize these

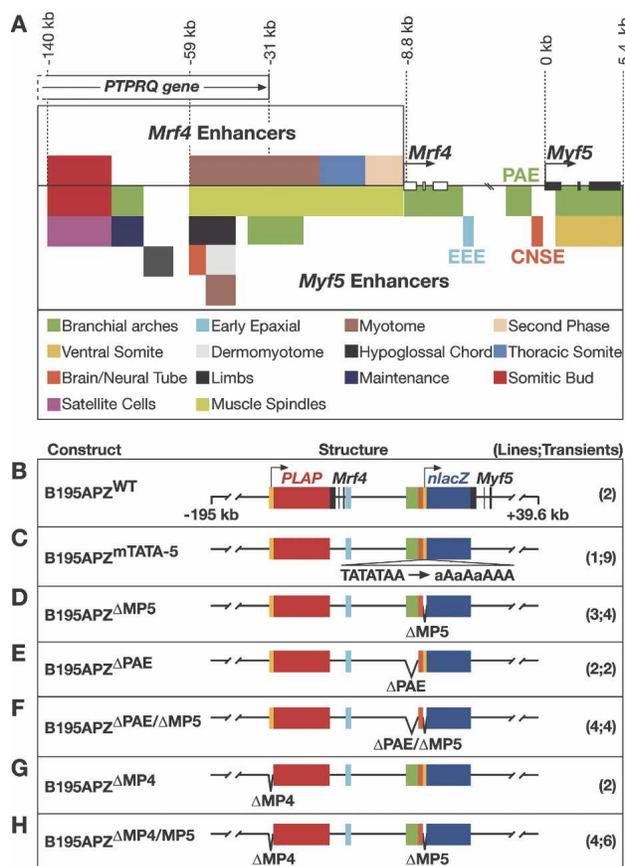


Figure 1. Maps of the regulatory elements in the *Mrf4/Myf5* locus and of the constructs generated for the study of global regulation. (A) Schematic representation of the regions involved in the transcriptional regulation of the *Mrf4/Myf5* locus. Each colored box represents transcription occurring at either different times or anatomical locations in embryonic or adult skeletal muscle. The color key is included below the map. (B) Schematic representation of construct B195APZ^{WT}, showing the location of the reporter genes (red box, human placental alkaline phosphatase gene [*PLAP*]; dark blue box, nuclear localized *lacZ* gene [*nlacZ*]); the EEE, PAE, and CNSE (pale blue, green, and orange boxes, respectively); the minimal promoters (yellow boxes); and the end points of the BAC clone. (C–H) Schematic representation of the different modifications introduced into B195APZ, and the number of independent transgenic lines and individual transient embryos analyzed (in brackets). (MP5) *Myf5* minimal promoter; (MP4) *Mrf4* minimal promoter.

signals into a single skeletal muscle fate. *Mrf4* is also differentially regulated in time and space, but the number of regulatory sequences involved is more limited (Carvajal et al. 2001). The majority of *Myf5* enhancers are upstream of *Mrf4* elements and closer to the *Mrf4* promoter than to that of *Myf5*, and most are located in introns of the adjacent *PTPRQ* gene (Fig. 1A), which is not expressed in skeletal muscle precursors. The locus thus provides an ideal system to investigate the mechanisms by which enhancer–promoter specificity is controlled so as to precisely regulate the rapidly changing and exquisitely precise expression patterns of these determination genes.

We propose the existence of a novel mechanism for the global transcriptional control of complex loci that depends on the maintenance of a series of equilibria between enhancers and promoters that are revealed by modification of the regulatory elements within the locus. Crucial to the establishment and fine tuning of the different equilibrium states are intergenic sequences that can initiate transcription when the locus is perturbed and can interact positively or negatively with the enhancers. Because of their activity, we term these elements transcription balancing sequences (TRABS).

Results

Deletion of the Myf5 minimal promoter allows some Myf5-specific elements to interact with the Mrf4 promoter

In *Drosophila* promoter–enhancer specificity can depend on the precise sequence of the TATA box (Ohtsuki et al. 1998; Butler and Kadonaga 2001). A putative TATA box is located between 183 and 177 base pairs (bp) upstream of the translational start site of *Myf5*. To identify the transcriptional start site we carried out 5'-rapid amplification of cDNA ends (RACE) amplification of total RNA from nontransgenic embryos and obtained a single amplification fragment. Multiple independent clones were sequenced that localized the transcriptional initiation site 155 bp upstream of the ATG, indicating that the putative TATA box is likely to be functional (data not shown).

We investigated whether the TATA box contributes to enhancer–promoter selectivity by mutating this element (TATATAA to aAaAaAA) in the context of BAC195APZ (constructs B195APZ^{wt}, B195APZ^{mTATA-5}) (Fig. 1B,C, respectively), which can fully recapitulate the expression patterns of *Mrf4* and *Myf5* (Carvajal et al. 2001). In B195APZ^{mTATA-5} embryos these patterns were identical to those in B195APZ^{wt} embryos (data not shown).

Since the TATA box is not required for correct enhancer–promoter specificity in this locus, we investigated the effects of deleting the entire *Myf5* minimal promoter (as functionally defined by Summerbell et al. 2000; see Supplemental Fig. 1) to generate B195APZ^{ΔMP5} (Fig. 1D). The timing and pattern of expression of *AP-Mrf4* is normal at 9.25 and 9.5 dpc (Fig. 2A,B) but then rapidly changes, incorporating aspects of

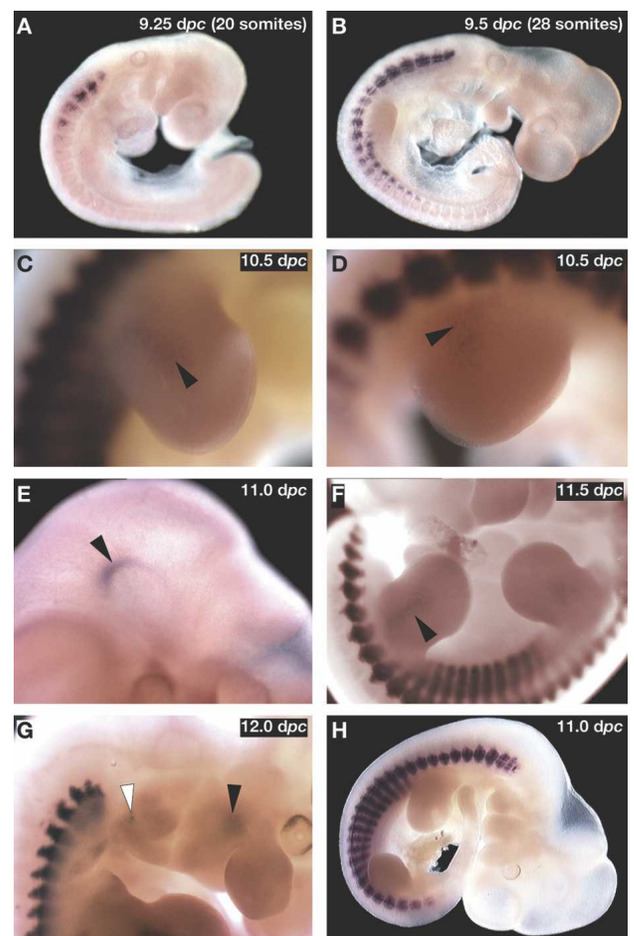


Figure 2. In the absence of the *Myf5* minimal promoter, some *Myf5*-specific enhancers drive *AP-Mrf4* expression. In the absence of the *Myf5* minimal promoter, expression of *AP-Mrf4* at 9.25 dpc (A) and 9.5 dpc (B) is indistinguishable from that of the wild type. At 10.5 dpc, *AP-Mrf4* is expressed in forelimbs (C, black arrowhead) and hindlimbs (D, black arrowhead). (E) At 11.0 dpc, *AP-Mrf4* is expressed in *Myf5*-specific domains such as the brain (black arrowhead). (F) By 11.5 dpc, *AP-Mrf4* expression is maintained in the limb buds, but is stronger in the hindlimb (black arrowhead). (G) By 12.0 dpc, *AP-Mrf4* expression is activated in derivatives of the mandibular and hyoid arches (black and white arrowheads, respectively). (H) In the control embryo, carrying construct B195APZ^{WT}, *AP-Mrf4* is not detected at these sites.

the *Myf5* pattern. At 10.5 dpc *AP-Mrf4* expression is detected in both fore- and hindlimbs (Fig. 2C,D, respectively), 48 h earlier than endogenous *Mrf4*. At 11.0 dpc *AP-Mrf4* is expressed in the brain (Fig. 2E), and is maintained in the limb buds and expressed weakly in the branchial arches (data not shown). The cells in the limbs and the arches in which this *AP-Mrf4* expression is seen do not, at these times, express *myogenin* (Yee and Rigby 1993) and thus have not entered the terminal differentiation program. By 11.5 dpc, expression is still visible in the limb buds (Fig. 2F), and at 12.5 dpc (Fig. 2G), and later (data not shown), weak *AP-Mrf4* expression is clearly seen in derivatives of the mandibular and hyoid arches,

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which are normally *Myf5* but not *Mrf4* expression domains. Thus, in the absence of the *Myf5* minimal promoter, *Myf5*-specific elements are able to activate the remaining promoter, adding *Myf5* expression domains to the normal pattern of *Mrf4* (which is shown in Fig. 2H; also shown in Fig. 6 of Carvajal et al. 2001). It thus might appear that straightforward promoter competition operates for some enhancers; e.g., those functioning in the limbs, brain, and branchial arches. Their productive engagement is always with the *Myf5* promoter but in its absence they engage with the *Mrf4* promoter to direct transcription that is temporally and spatially correct for *Myf5*. However, it is particularly interesting to note that the early epaxial enhancer (EEE), which drives the earliest expression of *Myf5* in the dorsomedial lip of the somite (Summerbell et al. 2000; Teboul et al. 2002), is not able to engage the *Mrf4* promoter even in this context (Fig. 2A,B), indicating that enhancer–promoter specificity in this locus is controlled by a different mechanism.

Myf5 expression is not abolished in the absence of its minimal promoter

To confirm that deletion of the minimal promoter abolishes *nlacZ-Myf5* expression, we analyzed B195APZ^{ΔMP5} embryos for β-galactosidase activity. Expression is completely abolished at 8.5 dpc but, surprisingly, weak *Myf5* expression is observed by 9.5 dpc in branchial arches and the caudal edges of the somites (Fig. 3A). By 10.5 dpc the intensity of expression is greatly increased and it is readily visible in arches, limb buds, and both rostral and caudal somitic edges, while still absent from the dorsomedial lip of the somite, the myotome, and the ventral somitic bud (Fig. 3B). At later stages, the expression pattern becomes virtually indistinguishable from that driven by the wild-type B195APZ (see Carvajal et al. 2001), except for the dorsomedial lip (Fig. 3C, 11.5 dpc) and the tail somites (Fig. 3D, 12.5 dpc) where it is missing. By 13.5 dpc the only difference from wild type is the missing expression in tail somites (Fig. 3E). Finally, from 14.5 dpc expression down-regulates (Fig. 3F), and it is abolished in all sites of normal *Myf5* expression in adult muscle (data not shown).

The unexpected fact that *Myf5* is widely expressed in the absence of its functionally defined minimal promoter means that the locus must contain at least one other, cryptic, promoter, and that many of the enhancers can activate it. This raises the question of whether the enhancers can engage with such sequences even when the minimal promoter is present. Moreover, it is clear that some of the enhancers that can activate the cryptic promoter—e.g., arch and limb enhancers—also activate the *Mrf4* promoter in the same cell population at the same time, and are thus not making a simple competitive choice. We suggest that they participate in an equilibrium system in which they can engage with multiple other sequences within a short time period. It is again noteworthy that the EEE cannot activate the cryptic promoters in this context, nor can the element that normally maintains *Myf5* expression after 14.5 dpc.

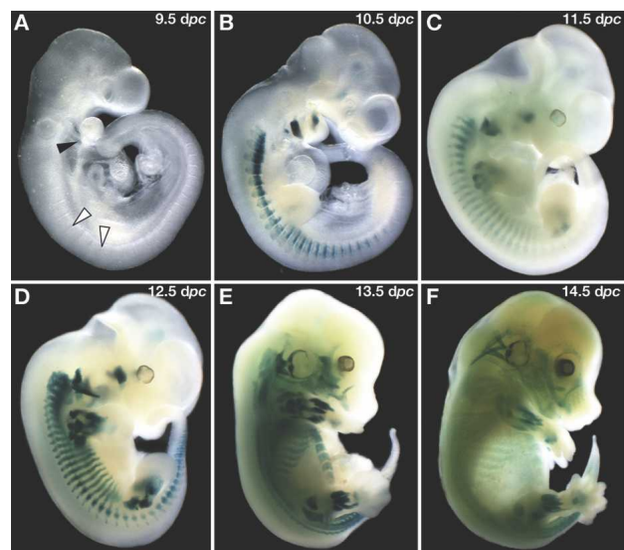


Figure 3. *Myf5* expression is not abolished in the absence of its minimal promoter. (A) *nlacZ-Myf5* expression can be detected at 9.5 dpc in the hyoid arch and the caudal edges of interlimb somites (black and white arrowheads, respectively). (B) By 10.5 dpc, expression is up-regulated in the dermomyotome of cervical somites, the caudal lips of interlimb somites, and branchial arches. (C) By 11.5 dpc, expression is clearly visible in fore- and hindlimbs, and myotomal expression is up-regulated for the first time. By 12.5 dpc (D), expression increases in all premuscle masses, is maintained at 13.5 dpc (E), and then starts to down-regulate at 14.5 dpc (F).

Modification or deletion of the *Myf5* minimal promoter activates novel initiation sites in the locus

To map the cryptic promoter(s) we carried out 5'-RACE on mRNA extracted from 13.5-dpc B195APZ^{wt}, B195APZ^{mTATA-5}, and B195APZ^{ΔMP5} embryos. The TATA-box mutation alters the initiation site, leading to a variety of transcripts with 5'-ends ranging from –650 to –100 bp relative to the translational start site, and very little, if any, initiation at the normal site (Fig. 4A [lane 3], B; Supplemental Fig. 1), indicating that this consensus TATA box is responsible for specifying the transcriptional start site of *Myf5* and that, under normal developmental conditions, its absence does not affect the expression pattern, while resulting in incorrect initiation. It also follows that transcription initiated at upstream start sites is correctly regulated by, as far as we can tell, all of the enhancers. For B195APZ^{ΔMP5} the *nlacZ*-specific nested primer set generates a ladder with fragments ranging from 50 to 1200 bp (Fig. 4A [lane 4], B; Supplemental Fig. 1), indicating that in the absence of the minimal promoter several transcripts are generated, starting within previously defined elements in the intergenic region: the proximal arch element (PAE), which in isolation is able to drive arch expression from 9.5 dpc to 12.5 dpc, and the CNS element (CNSE), which drives expression in the neural tube (Summerbell et al. 2000). We confirmed this using a different 5'-RACE protocol, which generated similar results (TaKaRa protocol) (data

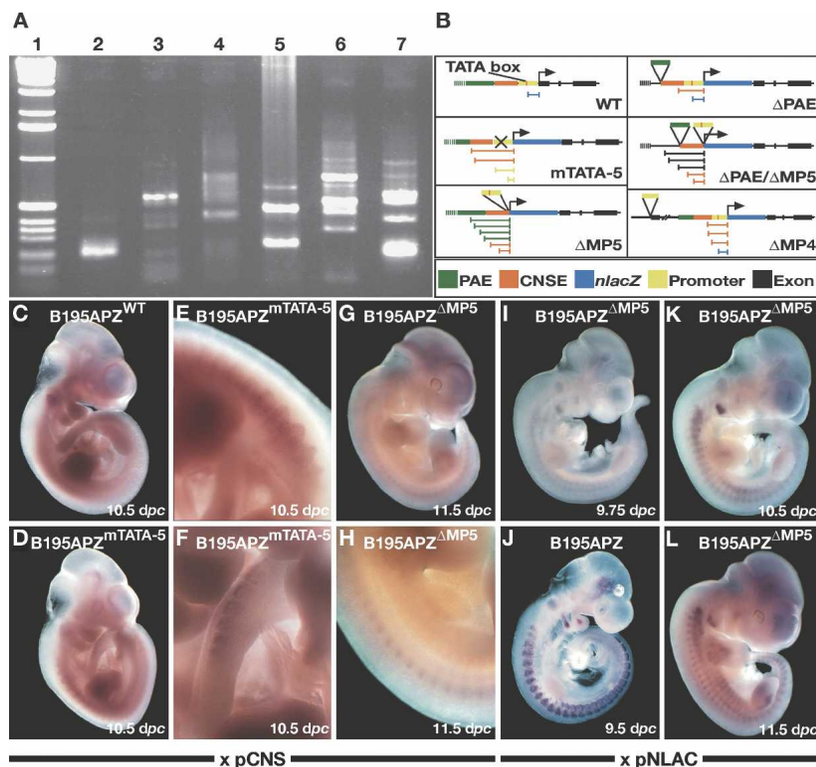


Figure 4. Alterations in the composition of the locus result in modifications of the transcriptional start site of *Myf5* as detected by 5'-RACE and in situ hybridization. The transcriptional start sites for the transgene change depending on the promoter/enhancer composition of the locus. Mutation of the TATA box in the *Myf5* promoter results in the activation of multiple transcription initiation sites. (A) Deletion of the *Myf5* minimal promoter, which activates a cryptic promoter, also results in multiple transcriptional starts, while deletion of the PAE or the *Mrf4* promoter has a strong influence on transcription initiation at *Myf5*, resulting in the activation of additional cryptic sites. (Lane 1) One-kilobase DNA marker (GIBCO-BRL). (Lane 2) Wild type. (Lane 3) B195APZ^{mTATA-5}. (Lane 4) B195APZ^{ΔAMP5}. (Lane 5) B195APZ^{ΔPAE}. (Lane 6) B195APZ^{ΔPAE/ΔAMP5}. (Lane 7) B195APZ^{ΔAMP4}. These putative initiation sites are represented in B, and have been shown to map to different enhancer elements upstream of the *Myf5* promoter. These transcripts can be detected by using riboprobes against noncoding sequences immediately upstream of the promoter (D–H), but not in B195APZ^{wt} embryos (C). Hybridization with a *nlacZ*-specific riboprobe shows low overall levels of transgene transcription (I,K,L) compared with a transgenic line carrying the wild-type BAC (J).

not shown). Because of the nature of 5'-RACE protocols we cannot rule out that some start sites are even further upstream.

To confirm the existence of these new transcripts we performed whole-mount in situ hybridization using a probe corresponding to the CNSE. Transcripts are not detected in nontransgenic (data not shown) or B195APZ^{wt} embryos (Fig. 4C), but transcripts containing this upstream sequence are detected in 10.5-dpc B195APZ^{mTATA-5} embryos (Fig. 4D), in the somites and limbs and very faintly in the branchial arches (Fig. 4E,F). Transcripts containing CNSE sequences are also detected in 10.5- and 11.5-dpc B195APZ^{ΔAMP5} embryos. At 10.5 dpc these transcripts are detected at low intensity in hyoid arch, somites, and limb buds (data not shown), while at 11.5 dpc transcripts are detected mainly in the dorsal somites (Fig. 4G,H). The low intensity of the signals indicates that deletion of the promoter reduces the levels of *Myf5* transcription. Indeed, hybridization of B195APZ^{ΔAMP5} embryos using an *nlacZ*-specific probe (pNLAC) shows that there is a significant reduction in transcription levels in all muscle precursors in early embryos (Fig. 4I) when compared with transgene expression in B195APZ^{wt} lines (Fig. 4J). At later stages (10.5 and 11.5 dpc) (Fig. 4K,L, respectively), there is an increase in *nlacZ-Myf5* expression levels, particularly in derivatives of the hyoid arch, recapitulating some of the differences in intensity observed with β -galactosidase staining of these lines (Fig. 3B). The novel transcripts are found only where *Myf5* is normally expressed; we saw no evidence for inappropriate expression.

The Mrf4 promoter is a key element in the maintenance of equilibria in the locus

We next generated a construct in which the *Mrf4* minimal promoter (as functionally defined by Black et al. 1995; see Supplemental Fig. 2) was deleted (B195APZ^{ΔMP4}) (Fig. 1G). The *Myf5* expression pattern in such embryos (Fig. 5A) is indistinguishable from that of B195APZ^{wt} embryos (Fig. 6A), while *AP-Mrf4* expression is completely abolished (data not shown), suggesting that all the cryptic promoters in the locus are located in the intergenic region. Because in the absence of the *Myf5* minimal promoter some of the *Myf5*-specific enhancers are able to drive both *nlacZ-Myf5* and *AP-Mrf4* expression, we generated construct B195APZ^{ΔMP4/ΔAMP5} (Fig. 1H), in which both promoters are absent, to ascertain if the *nlacZ-Myf5* expression was a result of read-through from the *Mrf4* promoter. At 9.5 dpc, construct B195APZ^{ΔMP4/ΔAMP5} drives *nlacZ-Myf5* expression in the dermomyotome and branchial arches but fails to drive dorsomedial lip or myotomal expression (Fig. 5B), showing that in the absence of the *Myf5* minimal promoter *nlacZ-Myf5* does not initiate at the *Mrf4* promoter. More importantly, comparison of the *nlacZ-Myf5* expression patterns of B195APZ^{ΔMP4/ΔAMP5} and B195APZ^{ΔAMP5} in the somites (Fig. 5B,C, respectively) shows that the *Mrf4* promoter prevents a somitic enhancer from activating the cryptic promoters at this time, and only by removing the *Mrf4* promoter is the enhancer able to drive *nlacZ-Myf5* in the dermomyotome. We infer that the enhancer preferentially interacts nonproductively with the *Mrf4*

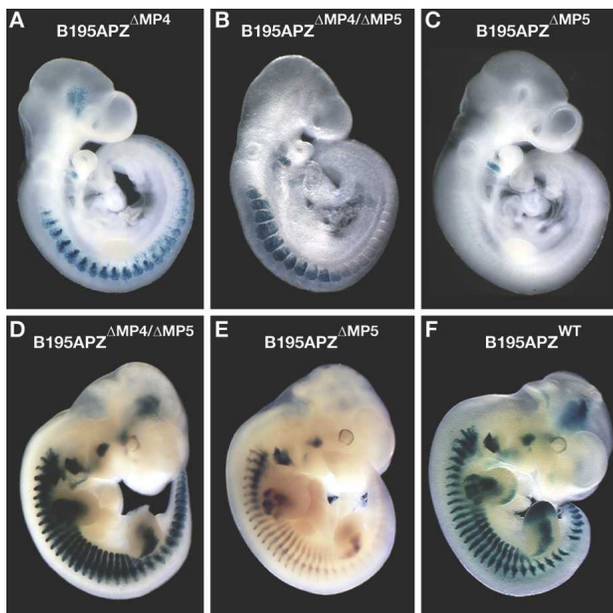


Figure 5. The *Mrf4* promoter is a key player in the enhancer-promoter equilibria in the locus. Deletion of the *Mrf4* minimal promoter does not affect the *nlacZ-Myf5* expression pattern (A), while the double MP4/MP5 deletion partially rescues the early (9.5 dpc) *Myf5* expression pattern (B), lost in the single MP5 deletion (C). The double deletion also rescues the expression pattern at 11.5 dpc, resulting in up-regulation of the transgene in all premuscle masses (D), while the construct carrying the single MP5 deletion is unable to drive full expression in limbs, interlimb, or tail somites (E). (F) The $B195APZ^{\Delta MP4/\Delta MP5}$ expression pattern is indistinguishable from that of $B195APZ^{WT}$ at this stage.

promoter when it is present. There is no expression in the dorsomedial lip, confirming that even if the *Myf5* minimal promoter is not present the EEE is sequestered by other sequences and prevented from activating the *Mrf4* promoter (see model in Fig. 7B). At later stages the double deletion results in up-regulation of *nlacZ-Myf5* in all muscle precursors (Fig. 5D) when compared with $B195APZ^{\Delta MP5}$ (Fig. 5E), and the pattern became indistinguishable from that driven by $B195APZ^{WT}$ (Fig. 5F). This shows that most, if not all, of the enhancers can activate the cryptic promoters if the *Mrf4* promoter is absent. Surprisingly, this deletion alters the *Myf5* initiation site, activating a site within the CNSE, and minor sites within the PAE in addition to the normal site (Fig. 4A [lane 7], B). The finding that deletion of the *Mrf4* promoter affects transcriptional initiation at *Myf5* indicates that both promoters are part of an equilibrium operating in the locus.

*The PAE is in equilibrium with other enhancers in the locus and with the *Mrf4* minimal promoter*

Expression in the branchial arches is subject to probably the most complex regulatory system in the locus, with at least five enhancers acting in the arches and their derivatives (Summerbell et al. 2000; Carvajal et al. 2001). Fur-

thermore, we showed that in the absence of the *Myf5* minimal promoter, *nlacZ-Myf5* expression in the hyoid arch is maintained or up-regulated at the transcriptional level, suggesting that the PAE strongly activates the cryptic promoters in the locus. To investigate this and the contribution of the PAE to the normal regulation of *Myf5*, we analyzed *nlacZ-Myf5* expression in $B195APZ^{\Delta PAE}$ embryos, in which the PAE had been deleted (Fig. 1E; Supplemental Fig. 1). Early *Myf5* arch expression was normal, but was down-regulated from 9.5 dpc (data not shown), although not completely abolished. We could not detect any other effects at any of the stages analyzed (9.5–13.5 dpc) (data not shown), indicating that sequences within the PAE are not required to

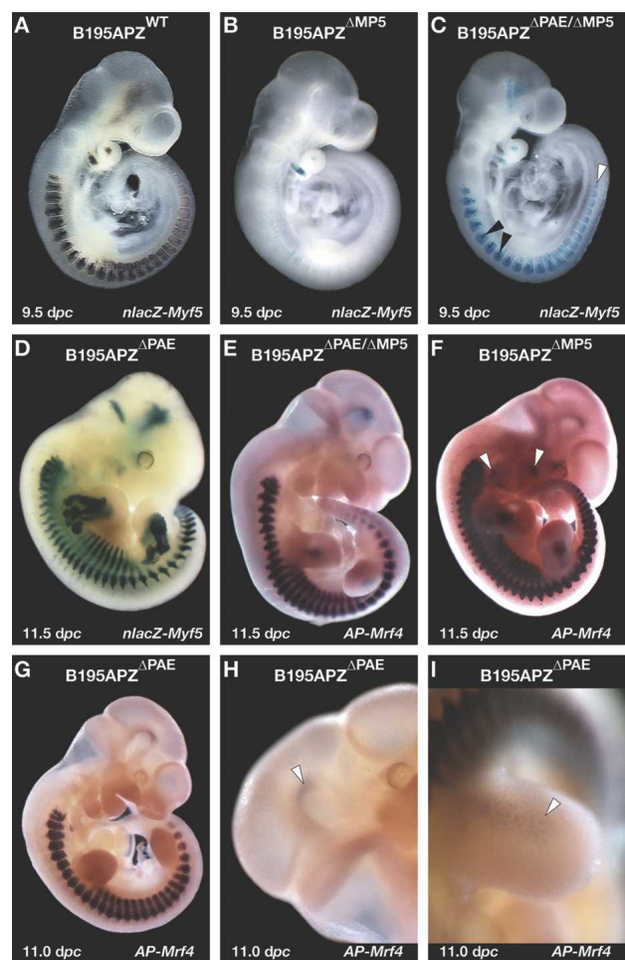


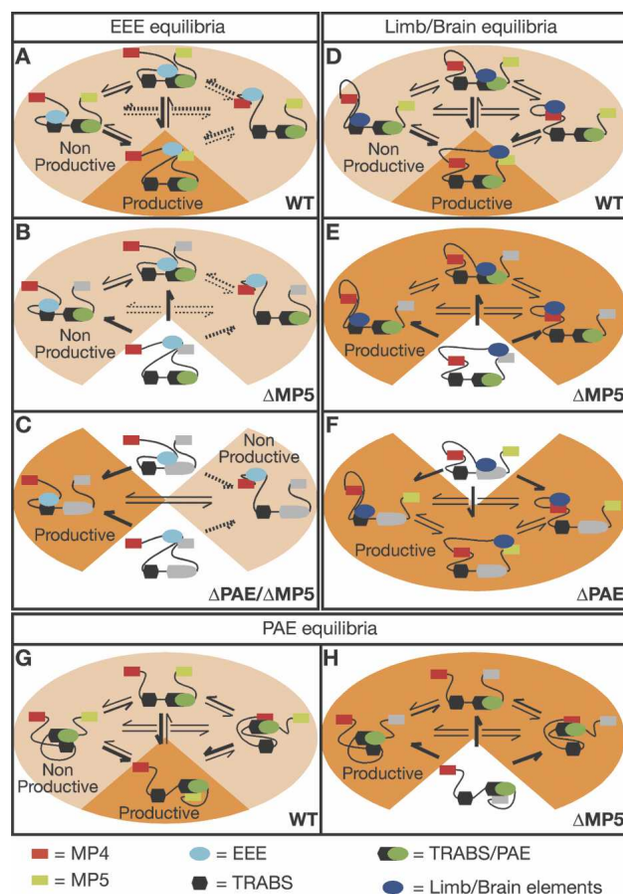
Figure 6. The PAE is essential for the maintenance of the equilibrium between enhancers and promoters in the locus. *nlacZ-Myf5* expression patterns driven by $B195APZ^{WT}$ (A) and $B195APZ^{\Delta MP5}$ (B) constructs at 9.5 dpc. (C) Deletion of both the *Myf5* minimal promoter and the PAE results in the rescue of the complete pattern of expression at this stage. (D) Single PAE deletion results in the loss of arch expression at 11.5 dpc, without any other effects on *Myf5* expression. Double PAE/MP5 deletion abolishes *AP-Mrf4* expression in the arches (E), which is readily detected in the single MP5 deletion (F). Single PAE deletion alters the expression of *Mrf4* (G) by activating *Myf5*-specific expression in the brain (H) and early in the limbs (I).

activate *Myf5* transcription at any other location. Surprisingly, deletion of the PAE alone results in the activation of an upstream start site, while the normal site downstream from the TATA box remains active (Fig. 4A [lane 5], B). The changes in the transcriptional start site of *Myf5* in the absence of the PAE indicate that it has dual functions, to drive late arch expression and to ensure correct transcriptional initiation at the minimal promoter.

Our data suggest that sequences within the PAE may contribute to the regulation of the equilibria within the locus, and we showed that the EEE is subject to particularly stringent regulation. To further explore the role of the PAE we analyzed B195APZ^{ΔPAE/ΔMP5} animals, lacking both the *Myf5* minimal promoter and the PAE (Fig. 1F). To our surprise, the full *Myf5* expression pattern was restored at 9.5 dpc (Fig. 6C), including expression in the dorsomedial lip of the somite, driven by the EEE (Summerbell et al. 2000; Teboul et al. 2002). The pattern was indistinguishable from that of B195APZ^{wt} (Fig. 6A), showing that the enhancers accurately regulate the cryptic promoters. Deletion of both the PAE and the *Myf5* minimal promoter allows the EEE to activate the other cryptic promoters to drive correct dorsomedial lip *Myf5* expression. The simplest explanation of these data is that in the absence of the *Myf5* minimal promoter the EEE interacts with sequences within the PAE interval in a way that prevents it from activating the cryptic promoters (see model in Fig. 7A–C).

By 11.5 dpc, *Myf5* arch expression is down-regulated in B195APZ^{ΔPAE/ΔMP5} embryos (data not shown), reproducing the timing of arch expression driven by construct B195APZ^{ΔPAE} (Fig. 6D). Because the down-regulation of *Myf5* arch expression in B195APZ^{ΔPAE} overlaps in time with the up-regulation of *Mrf4* expression in B195APZ^{ΔMP5} lines, we hypothesized that in the absence of the *Myf5* minimal promoter the PAE engages with the *Mrf4* minimal promoter to drive this expression. Analyses of the *AP-Mrf4* expression pattern at 11.5 dpc in B195APZ^{ΔPAE/ΔMP5} embryos (Fig. 6E) show that this is indeed the case, as these embryos lack the arch expression seen in B195APZ^{ΔMP5} embryos (Fig. 6F), while the remainder of the pattern was identical (data not shown). In the absence of both the PAE and the minimal promoter, transcription is still initiated within the CNSE and at locations upstream of the deleted PAE (Fig. 4A [lane 6], B), indicating that a variety of sequences in the locus are able to initiate transcription in the absence of the minimal promoter.

We showed that the PAE has a dual function, controlling the expression of *Myf5* in the branchial arches and ensuring correct transcriptional initiation at the *Myf5* promoter. In the absence of the *Myf5* minimal promoter the PAE is involved in driving expression of *AP-Mrf4* in



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the arches while double deletion of the PAE and the *Myf5* minimal promoter reveals that sequences within the PAE interval sequester some *Myf5*-specific enhancers, thus preventing their interaction with other cryptic promoters in the locus. If the role of the PAE interval in the maintenance of the equilibria is crucial, deletion of this element in isolation should result in alteration of the enhancer–promoter specificity in the locus. While the changes in the pattern of *nlacZ-Myf5* are restricted to the absence of early arch expression (Fig. 6D), there are important changes to the *AP-Mrf4* pattern (Fig. 6G), which incorporates expression in the brain (Fig. 6H) and the limb buds (Fig. 6I). Therefore, in the absence of the PAE at least some enhancers are able to activate basal promoters that they cannot access when it is present, indicating that sequences within the PAE interval play a key role in regulating the overall enhancer–promoter equilibria (see model in Fig. 7D–H).

Discussion

Promoter–enhancer equilibria

Eukaryotes employ many different mechanisms to ensure that transcriptional units can respond appropriately to the surrounding environment in order to regulate the activity levels of the many proteins encoded in their genomes. Crucial to this response is the control of transcriptional regulators themselves, as downstream gene activation is mainly dependent on the correct control of transcription factor activity. In multicellular organisms complex regulatory mechanisms are required to ensure tight regulation of tissue-specific gene expression. These mechanisms must be even more complex when two or more genes, with distinct expression patterns, are linked in the genome, with the possibility of enhancer cross-talk even greater. Furthermore, it has been shown that gene duplication is usually accompanied by the sharing of enhancers belonging to the protogene, and the specialization of the duplicated genes for individual enhancers, usually resulting in nonoverlapping patterns that when combined cover the entire expression pattern of the original gene (Bruce et al. 2001; Locascio et al. 2002; de Souza et al. 2005). We propose that the regulation of such loci involves additional mechanisms to ensure the correct enhancer–promoter specificity.

In multigene loci, several mechanisms have been described to explain how promoter–enhancer specificity is established. These seem to rely on a hierarchical architecture in which promoters compete for the different enhancers, with sequences either within (TATA box, downstream promoter elements, initiator sequences) or outside (tethering elements) the promoters used to modify the strength of the competition, and/or block such interactions (silencers, insulators, boundary elements). Other elements involved are the LCRs, responsible for the initiation and/or maintenance of tissue-specific gene expression. The most widely studied LCR is that of the β -*globin* gene cluster, which has been shown to be required for high levels of β -*globin* expression (for reviews, see Fraser and Grosfeld 1998; Li et al. 2002).

Importantly, the individual *globin* genes compete for the LCR and, as with other forms of promoter competition, the LCR only interacts with one promoter at a time (Wijgerde et al. 1995). Finally, another *cis*-regulatory element, the promoter targeting sequence, has also been shown to contribute to promoter–enhancer specificity by selectively bypassing the effects of specific insulators in the *Drosophila Abd-B* gene (Zhou and Levine 1999).

By altering the regulatory element composition of the *Mrf4/Myf5* locus we now reveal a new mechanism. In the absence of the *Myf5* minimal promoter, some of the enhancers that interact with it in the normal context are recruited by the *Mrf4* promoter to drive *Myf5*-specific expression. Strikingly, other sequences in the locus can also recruit these enhancers and assemble the transcriptional machinery to drive *Myf5* expression in the absence of its promoter. Not all of the enhancers behave in the same manner, and thus, for example, expression is up-regulated in the hyoid arch at 9.5 dpc and myotomal expression is down-regulated at early stages, while dorsomedial lip expression mediated by the EEE is completely abolished. Because some of the enhancers drive expression of both *AP-Mrf4* and *nlacZ-Myf5* in the same location and at the same stage, this novel mechanism does not correspond to any previously described on/off-switch model. Deletion of the *Mrf4* minimal promoter alters the expression of *Myf5* at two time points during development. Early *Myf5* expression is missing in the somitic bud, and late expression is abolished throughout the embryo and in the adult. Even more intriguingly, the deletion of both the PAE and the *Myf5* minimal promoter results in pattern reversion of *Myf5* expression to that of the wild type. Analyses at the RNA level indicate that deletion of either promoter results in modifications to the transcriptional start site for *nlacZ-Myf5*, further evidence of the *cis*-regulatory interactions between the two promoters in the locus. But these alterations in transcription initiation are not restricted to constructs missing the natural promoters. Indeed, deletion of the PAE also results in alteration of transcriptional initiation and changes in the pattern of *Mrf4* expression. These observations indicate that the locus is not regulated by straightforward promoter competition, and that elements different from the core promoters are involved in enhancer–promoter selectivity, elements that we name TRABS.

Cryptic promoters or TRABS

The term cryptic promoter is commonly used in the literature to refer to a fragment of genomic DNA that is able to act as a promoter in certain circumstances but does not normally act. In this way, in several chromosomal translocations involving the *c-myc* locus, and giving rise to mouse plasmacytomas, it has been shown that the loss of the endogenous *c-myc* promoter in the derivative chromosome does not imply the silencing of *c-myc*, and that the generation of a truncated *c-myc* protein is due to the activation of bipolar cryptic promoters in the locus (Calabi and Neuberger 1985; Murphy et al. 1986). A

cryptic promoter located 2 kb upstream of the *Slp* gene in the *eIF4G* locus, and originally defined as an internal ribosomal entry site (IRES), has been recently shown to be able to increase by 850 times the expression of an associated *Renilla luciferase* gene (Han and Zhang 2002). The authors also identified additional cryptic promoters in the 5'-untranslated region (UTR) of the *Bad* and *Sno* genes, previously defined as IRES, and concluded that cryptic promoter activities in the 5'-UTR sequences of cellular mRNAs are more prevalent than anticipated.

Deletion of the *Myf5* minimal promoter in the context of B195APZ results in the activation of a cryptic promoter(s) in the locus. Furthermore, we showed that deletion of a single enhancer (the PAE) is enough to activate a cryptic promoter, as revealed by 5'-RACE analyses. It is not clear at the moment if these sequences are one or many, although the variation in the transcriptional start site for the *nIacZ-Myf5* transgene in the different deletion constructs indicates that multiple sequences in the locus may be activated as cryptic promoters. Furthermore, we showed that the cryptic promoters are able to engage with most of the *Myf5*-specific enhancers in a context-dependent manner, changing their specificity according to the promoters in the locus, suggesting that the different sequences are in competition with the natural promoters and with each other. We propose that many of the sequences described as cryptic promoters in the literature are in fact TRABS, and that their function is to establish an equilibrium with the promoter(s) in the locus to ensure the fine control of the interactions between enhancers and natural promoters required for the correct transcriptional initiation of gene expression.

Analysis of the equilibrium

Deletion of the *Myf5* minimal promoter leads to *Mrf4* expression in a number of sites (brain, branchial arches, and limbs) that are normally *Myf5*-specific, which might be taken as indicating that for some enhancers there is straightforward promoter competition. However, other enhancers, most notably the EEE, cannot engage with the *Mrf4* promoter even when the *Myf5* promoter is absent, indicating that a different mechanism must operate. Many of the enhancers can thus engage with the cryptic promoters in an entirely specific fashion, but their ability to do so is modulated not only by the *Myf5* promoter, which would be expected, but also by the *Mrf4* promoter and the PAE. A key feature of the model that we propose (Fig. 7) is that these interactions between the enhancers and the cryptic promoters occur normally (Fig. 7A,D,G), and that they dynamically regulate the choices made as to which enhancers should engage with which promoter, in which progenitor cell population and at what time.

The second key observation is that in the absence of the *Myf5* minimal promoter, enhancers operating in the arches (Fig. 7H), the limbs or the brain (Fig. 7E), engage with both the *Mrf4* promoter and the cryptic promoters in the same progenitor cell population at the same time.

It is clear that the ways in which the enhancers engage with the promoters are not controlled by an on/off switch. Rather, we propose that the enhancers are continuously interacting with a number of transcription initiation sites in the locus, and which interaction leads to the productive initiation of transcription is dictated by the instructive signals that the progenitor cell is receiving.

The model is further supported by the deletion of the minimal promoters of both *Mrf4* and *Myf5*. While most of the early *Myf5* somitic pattern is abolished in the absence of the *Myf5* minimal promoter, deletion of the *Mrf4* promoter results in partial rescue of the somitic pattern. We suggest that the enhancer driving this expression cannot engage with the cryptic promoters because it is sequestered nonproductively by the *Mrf4* promoter. When the latter is removed, the enhancer is released and engages productively with the cryptic promoters. Similarly, the EEE cannot activate the cryptic promoters in this context because it is sequestered by sequences within the PAE interval. This is a third key feature of our model, the regulation of the locus depends on interactions between enhancers and promoters, both conventional and cryptic, that do not lead to the initiation of transcription.

It is striking that in our initial experiments the EEE was not seen to interact with anything other than the *Myf5* minimal promoter, implying that there must be a mechanism that prevents its engagement with the *Mrf4* and cryptic promoters (Fig. 7B). But deletion of the PAE and the *Myf5* minimal promoter allows the EEE to activate one or more cryptic promoters (Fig. 7C). We thus infer that the EEE normally engages productively only with the *Myf5* minimal promoter but that it is in an equilibrium between that promoter and the TRABS within the PAE interval. Only when both are removed is the EEE able to activate the cryptic promoters, although it is still, by an unknown mechanism, prevented from activating the *Mrf4* promoter. It is important to mention that it is not clear yet if the PAE interval contains two separable functional elements (the arch enhancer and the TRABS) or if they are interdigitated or even synonymous, and we are currently investigating this point. We suspect that if all the TRABS were removed, then the EEE would be able to activate the *Mrf4* promoter, and we are examining this issue.

The *Mrf4* 'knockout' and the *Myf5* 'knock-in' revisited

Three different *Mrf4* knockout alleles were generated more than a decade ago by the laboratories of Arnold, Olson, and Wold (for review, see Olson et al. 1996 and references therein). Although all three alleles abolished *Mrf4* expression, variable *cis*-acting effects on the linked *Myf5* gene were observed, resulting in homozygous phenotypes ranging from full viability to complete lethality. In a previous article (Olson et al. 1996), we and others hypothesized that the presence of the strong *pgk* promoter on the selectable marker was partially responsible for some of the *cis*-effects observed, but we did not have an explanation for the mechanism or for the apparent

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discrepancy in phenotypes between the two alleles where the construction of the targeted allele was very similar but the resulting phenotypes widely different. The Olson allele, in which the *pgk-neo*-selectable marker is transcribed in the opposite direction to *Mrf4* and *Myf5*, presented the mildest of phenotypes, with some rib malformations and no overt problems with myogenesis. The Arnold and Wold alleles carried the *pgk-neo*-selectable marker in the same orientation as *Mrf4* and *Myf5*, and while both presented myogenic defects, some animals carrying the Wold allele survived to adulthood while none of the animals carrying the Arnold mutation survived after birth.

The work presented here readily explains these results in that the presence of the strong *pgk* promoter would alter the transcriptional equilibria in the locus by competing with the endogenous promoters and TRABS, resulting in the misregulation of *Myf5*. Furthermore, in the Arnold allele, which resulted in a double *Mrf4/Myf5* knockout, 102 bp of the *Mrf4* minimal promoter were crucially removed, including the transcriptional start site (Braun and Arnold 1995). As we showed that the *Mrf4* minimal promoter is a major player in establishing/maintaining the equilibria in the locus and is essential to determine the correct transcriptional initiation site for *Myf5*, it is likely that even its partial removal compromises *Myf5* promoter function, as we showed with our B195APZ^{ΔMP4} lines, in which *Myf5* expression in the somitic bud, late embryonic phases and the adult is abolished. Thus, the combination of the partial promoter deletion and the inclusion of an additional promoter in the locus could have resulted in the complete down-regulation of the linked *Myf5* gene, resulting in the double knockout characteristic of this allele.

More recently, Kassar-Duchossoy et al. (2004) have shown other *cis*-effects in the locus in a series of *Myf5* knock-in lines. The original *Myf5^{nlacZ}* allele (Tajbakhsh et al. 1996a) and two new alleles, *Myf5^{GFP-P}* and *Myf5^{loxP}*, show differential effects on the levels of *Mrf4*. Expression of *Mrf4* is abolished at 10.5 dpc in *Myf5^{nlacZ/nlacZ}* embryos, and reduced to half of the normal levels in *Myf5^{nlacZ/+}* littermates. A similar knock-in allele, *Myf5^{GFP-P}* also shows alteration in *Mrf4* levels, in both heterozygous and homozygous animals, but this expression is not completely abolished. Finally, in *Myf5^{loxP/loxP}* embryos, *cis*-effects were not observed. Although the three alleles abolish functional *Myf5* protein and have similar genomic structures, the selectable markers and associated promoters used are different. In the *Myf5^{nlacZ}* allele, the neo cassette is driven by the RNA pol II promoter (Tajbakhsh et al. 1996b), while in the *Myf5^{GFP-P}* allele the puromycin cassette is driven by the *pgk* promoter (Kassar-Duchossoy et al. 2004). It is therefore possible to speculate that the phenotypes of these two alleles that introduce a foreign promoter into the locus are due to their ability to disrupt the delicate balance required to set the enhancer–promoter specificity. Differences in phenotypes between these two alleles would be caused by the different affinities of the foreign promoters for the existing transcriptional elements in

the locus. Furthermore, Kaul et al. (2000) have shown that the rib phenotype observed in two of the original *Myf5* knockout alleles is caused by the selectable marker, although in this case the mechanism is unclear.

Conclusion

There are now a number of examples of genes with regulatory elements located far from the promoter, and it is noteworthy that they tend to encode either signaling molecules (e.g., Shh) or transcription factors (e.g., Sox9) involved in the control of cell fate decisions (Sagai et al. 2005; Bien-Willner et al. 2007). The extremely complicated enhancer organization in the *Mrf4/Myf5* locus is also not unique. *Sox2* is expressed throughout the early nervous system, but this overall expression pattern is the summation of the activities of a number of dispersed enhancers, each of which operates in a particular population of progenitor cells at a particular time (Uchikawa et al. 2003). Structuring the regulatory elements in this way, together with the novel regulatory mechanism described here, would seem to be particularly appropriate for genes encoding transcription factors, like *Myf5* and *Sox2*, which determine cell fate and have to be induced widely in the embryo in response to a variety of inductive signals.

Our data lead to a model that is quite distinct from those developed from analyses of the *β-globin* and *Hox* loci. They imply that the enhancers operate in what might be thought of as a neural network model. They are continuously interacting with not only the two conventional promoters in the locus but also with a number of sequences that can act as cryptic promoters, which we call TRABS. These interactions must be modulated by the inductive signals that determine skeletal muscle identity and lead to the activation of the two genes in an exquisite temporal and spatial pattern involving many different progenitor cell populations at many stages of development.

Materials and methods

Homologous recombination in Escherichia coli

Details on the construction of modifying cassettes can be found in the Supplemental Material.

We used the linear recombination method (Swaminathan et al. 2001) with some modifications. BAC clones were transferred into DY380 cells (kindly provided by Neil Copeland, National Cancer Institute-Frederick, Frederick, Maryland), which carry the temperature-inducible λ -recombinase system.

Single DY380 colonies carrying the desired BAC clone were isolated and grown overnight at 32°C. Two-hundred-twenty microliters of these cultures were used to seed 11 mL of LB and were incubated at 32°C with vigorous shaking (>300 rpm) until OD₆₀₀ was 0.6 ± 0.05. Cultures were then transferred to a 42°C waterbath and incubated for 15 min. Cultures were transferred to ice, incubated for 20 min, made electrocompetent by standard methods, and resuspended in a final volume of 30 μ L.

Modifying cassettes (2 nmol for TATA box mutation; 1 μ g for other cassettes) were denatured, ethanol-precipitated, and re-

suspended in 20 μ L of ice-cold H₂O. The denatured cassettes were mixed with 30 μ L of electrocompetent DY380 cells carrying the BAC clone to be modified. Electroporations were carried out using the Bio-Rad GenePulser II system (1.75 kV, 100 Ω , 25 μ F). Electroporated cells were diluted 1:10⁶ into a final volume of 50 mL of LB and aliquoted into a single 96-well plate (500 μ L per well), resulting in an average of 20–50 cells per well. After overnight incubation at 32°C, pools were screened using the appropriate primers (see the Supplemental Material). Two to six positive pools were plated to obtain single colonies, and a second round of screening was carried out to identify single positives. Clones were sequenced using primers outside the homology arms.

Generation of transgenic mice

All in vivo experimentation was performed according to United Kingdom Home Office Regulations. BAC DNA was prepared using the QIAGEN Maxiprep kit (Qiagen) as described previously (Carvajal et al. 2001). After dialysis against microinjection buffer (10 mM Tris-HCl at pH 7.5, 0.1 mM EDTA at pH 8.0, 100 mM NaCl), DNA was diluted to 1.5 ng/mL in microinjection buffer and used to inject fertilized mouse eggs from CBA/Ca \times C57Bl/6 crosses, as described previously (Yee and Rigby 1993).

Histochemical staining

β -Galactosidase: Embryos were fixed overnight in Mirsky's fixative (National Diagnostics) at 4°C, washed three times in PBSA (Ca²⁺, Mg²⁺-free phosphate-buffered saline)/0.02% Nonidet P-40 for 20 min at room temperature, placed in 10 mL of X-gal solution [5 mM K₃Fe(CN)₆, 5 mM K₄Fe(CN)₆ · 3H₂O, 2 mM MgCl₂, 0.02% Nonidet P-40, 0.4 mg/mL X-Gal in PBSA] for 2–20 h (depending on stage) at 37°C, and post-fixed in Mirsky's fixative.

Alkaline phosphatase: Embryos were fixed in 4% paraformaldehyde in PBSA overnight at 4°C, rinsed twice in PBSA + 2 mM MgCl₂, and washed in PBSA + 2 mM MgCl₂ for 10 min. Endogenous phosphatases were inactivated by incubation for 1 h in PBSA + 2 mM MgCl₂ at 65°C. Embryos were equilibrated in AP-buffer (100 mM Tris-HCl at pH 9.5, 100 mM NaCl, 50 mM MgCl₂, 0.1% Tween 20). Embryos were washed once in ice-cold AP-buffer, transferred to AP-staining buffer (AP-buffer + 0.1 mg/mL of BCIP; 0.5 mg/mL NBT), and stained for 3–10 h (depending on stage) in the dark at 4°C. The reaction was stopped by washing in ice-cold PBSA containing 2 mM EDTA in the dark for 24 h at 4°C. Embryos were post-fixed in Mirsky's fixative adjusted to pH 5.0.

RACE

Embryos were obtained from timed matings, taking the day of vaginal plug as 0.5 dpc, and dissected in PBSA. For the identification of transgenic embryos, left forelimbs were transferred to single tubes containing 1 mL of X-gal solution (see above) and incubated at 37°C until coloration was detected. Single 13.5-dpc embryos were homogenized in 1 mL of TRI reagent (Sigma) using disposable plastic pestles and RNA isolated following the manufacturer's recommendations. RNA was resuspended in a final volume of 500 μ L of DEPC-treated H₂O. We used the SMART RACE Amplification Kit (BD Biosciences) following manufacturer's recommendations. Single-stranded cDNA was amplified by PCR using the Advantage 2 PCR Kit (BD Biosciences) using either *Myf5*- or *nIacZ*-specific primers (Supplemental Material). Single amplification bands were gel-purified and cloned into pCR-TOPO (Invitrogen) for sequencing.

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