c-Kit triggers dual phosphorylations, which couple activation and degradation of the essential melanocyte factor Mi

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Microphthalmia (Mi) is a bHLHZip transcription factor that is essential for melanocyte development and postnatal function. It is thought to regulate both differentiated features of melanocytes such as pigmentation as well as proliferation/survival, based on phenotypes of mutant mouse alleles. Mi activity is controlled by at least two signaling pathways. Melanocyte-stimulating hormone (MSH) promotes transcription of the Mi gene through cAMP elevation, resulting in sustained Mi up-regulation over many hours. c-Kit signaling up-regulates Mi function through MAP kinase phosphorylation of Mi, thereby recruiting the p300 transcriptional coactivator. The current study reveals that c-Kit signaling triggers two phosphorylation events on Mi, which up-regulate transactivation potential yet simultaneously target Mi for ubiquitin-dependent proteolysis. The specific activation/degradation signals derive from MAPK/ERK targeting of serine 73, whereas serine 409 serves as a substrate for p90 Rsk-1. An unphosphorylatable double mutant at these two residues is at once profoundly stable and transcriptionally inert. These c-Kit-induced phosphorylations couple transactivation to proteasome-mediated degradation. c-Kit signaling thus triggers short-lived Mi activation and net Mi degradation, in contrast to the profoundly increased Mi expression after MSH signaling, potentially explaining the functional diversity of this transcription factor in regulating proliferation, survival, and differentiation in melanocytes.

[Key Words: Microphthalmia, c-Kit, steel factor, MAPK, p90 Rsk, ubiquitin]
survival. The role of Mi in pigmentation has emerged from evidence that its expression is potently induced by the pigmenting factor melanocyte-stimulating hormone (MSH) [Bertolotto et al. 1998a; Price et al. 1998b] as well as recognition that Mi can up-regulate transcription through a conserved consensus element in the promoters of the major pigment enzyme genes [Bentley et al. 1994; Hemesath et al. 1994; Yasumoto et al. 1994]. Evidence of a major role for Mi in survival/proliferation of melanocytes comes from the mi<sup>v<sup>1<sup> mouse mutation that displays nearly normal neonatal melanocyte numbers followed by precocious melanocyte loss over several months of age (premature gray/white) [Lerner et al. 1986]. This phenotype is consistent with an essential role for Mi in post-developmental melanocyte proliferation or survival. In addition, the position of Mi downstream of Steel/c-Kit signaling is consistent with mitogenic or survival signals known to be stimulated by Steel/c-Kit in a variety of contexts [Andrews et al. 1994; Hassan and Zander 1996; Sykora et al. 1997]. Mi has also been shown to regulate c-Kit expression transcriptionally in mast cells [Tsujimura et al. 1996], suggesting the possibility of homeostatic regulation among these factors.

As a target of at least two signaling pathways, MSH and c-Kit, Mi may reside at a pivotal position for its ability to trigger alternative transcriptional programs. Although much remains to be learned about the spectrum of genes activated by Mi in melanocytes, it is plausible that different genes are targeted in distinct contexts and that the transcriptional activity of Mi may, therefore, be tightly regulated in a signal-dependent fashion.

Both the MSH and c-Kit signaling pathways up-regulate the transcriptional activity of Mi, but they do so in very different ways. MSH stimulation up-regulates cAMP and stimulates new transcription of Mi through a cAMP response element (CRE) in the Mi promoter in melanocytes. Thus, MSH stimulation profoundly increases Mi protein expression over the course of hours [Bertolotto et al. 1998a; Price et al. 1998b]. In contrast, c-Kit stimulation produces very rapid MAPK-mediated phosphorylation of Mi, producing enhanced recruitment of p300/CREB-binding protein [CREB-binding protein] [Price et al. 1998a], the coactivator family that interacts with and modulates the transcriptional activity of Mi [Sato et al. 1997]—all occurring over the course of minutes. The kinetic differences between these alternative means of up-regulating Mi are significant and could contribute to the different biological consequences of stimulating these signaling pathways. In other settings, transcriptional activity has been suggested to rely on proteolytic degradation of nuclear receptors [Nawaz et al. 1999], suggesting such phenomena could be of widespread importance.

In the current study we investigated the consequences of c-Kit signaling on Mi stability and function. We show that Mi is targeted for rapid ubiquitin-dependent proteolysis with Steel factor stimulation. The specific signals were found to be phosphorylation by either MAPK at serine 73 or Rsk-1 at serine 409. Double serine-to-alanine mutations at these two residues result in a protein that is both profoundly stable and transcriptionally inactive. These c-Kit-induced phosphorylations thus produce coupled, short-lived activation-destruction signals on the nuclear target Mi.

**Results**

**Mi is degraded after c-Kit stimulation**

The observation that Mi protein is degraded after c-Kit signaling was first made in the study of Sl stimulation of human melanoma cells. Western blots using a Mi-specific monoclonal antibody showed that Sl stimulation produced an initial mobility shift of Mi due to MAPK/ERK phosphorylation on serine 73 as previously described [Hemesath et al. 1998]. Subsequent to this shift, Mi protein levels appeared to diminish over time (Fig. 1A, left). Previous studies demonstrated that Steel/c-Kit signals are transmitted to Mi through MAPK/ERK [Hemesath et al. 1998]. To test whether this same pathway was triggering Mi degradation, the MAPK/ERK pathway inhibitor PD98059 was used (Fig. 1A) and was found to prevent both the c-Kit-dependent mobility shift and c-Kit-induced Mi degradation. This observation suggested that phosphorylation of ser-73 or another MAPK-dependent signal was responsible for Mi degradation. To clarify the fate of Mi protein after Sl signaling, de novo protein synthesis was inhibited by incubation with cycloheximide and the kinetics of Mi degradation were analyzed. With cycloheximide, Sl produced the same MAPK-dependent mobility shift in Mi, followed by more pronounced degradation of Mi (Fig. 1B, left). There was no reappearance of the lower band with arresting protein synthesis in these cells (Fig. 1B, left). The drug PD98059 again stabilized Mi protein levels over time (Fig. 1B, right). Pulse/chase analysis showed the same changes in Mi stability as cycloheximide decay kinetics (data not shown).

**A**

| Sl (hr) | -MEK Inhibitor | MEK Inhibitor |
|--------|----------------|--------------|
| 0      | 0.5            | 3            |
| 0      | 0.5            | 3            |
| 0      | 0.5            | 3            |
| 0      | 0.5            | 3            |

**B**

| Sl-Cl1 (hr) | MEK Inhibitor | MAPK Inhibitor |
|-------------|---------------|----------------|
| 0           | 0.5           | 3              |
| 0           | 0.5           | 3              |
| 0           | 0.5           | 3              |

**Figure 1.** c-Kit activation produces MAPK-dependent Mi degradation. [(A)] Human melanoma cells [501 Mel] were stimulated at the c-Kit receptor with recombinant Steel factor [Sl] in the presence or absence of the MEK inhibitor drug PD98059 and whole cell extracts probed with monoclonal antibody against Mi (doublet) or α-tubulin-loading control at the time points indicated. [(B)] c-Kit stimulation was also carried out in the presence of cycloheximide [CHX] to reveal Mi degradation kinetics.
Mi degradation is proteasome dependent

Mi from melanoma cells (501 Mel or MeWo) and melanocytes was examined to determine whether Mi instability involved proteasome-mediated degradation. Proteasome inhibition using MG132 or lactacystin [Fig. 2A–E] prevented Sl/c-Kit-induced Mi degradation in melanoma cells as well as primary human melanocytes [Fig 2A–C]. The proteasome inhibitor MG115 showed the same effect (data not shown). Inhibition of Mi degradation traps Mi in the upper phosphorylated form. Over time there is appearance of the lower band suggesting a dephosphorylation event. Treatment of cells with okadaic acid, a phosphatase 1 and 2A inhibitor, prevented the appearance of the lower migrating Mi isoform after 2 hr of Sl stimulation [Fig. 2C, right], suggesting that phospho-Mi may be subject to both degradation and dephosphorylation events in these cells. The same results were found with either murine melanoma cells [Fig. 2D] or murine melanocytes [Fig. 2E]. For murine cells, MAPK stimulation was triggered with phorbol ester (TPA) in lieu of Sl because these cells lack functional c-Kit receptor. The results from multiple cell systems suggest that Sl/c-Kit- and MAPK-dependent destabilization of Mi involves targeting of Mi to the proteasome degradation pathway.

c-Kit stimulation triggers ubiquitination of Mi

Proteins that are degraded by the proteasome machinery in cells are first marked for this fate by the covalent addition of polyubiquitin chains to specific lysine residues [Chau et al. 1989; Goldberg and Rock 1992; Hershko and Ciechanover 1992; Mahaffey et al. 1993; Palombella et al. 1994; Goldberg 1995]. To assess whether Mi undergoes covalent ubiquitination, two types of experiments were done. In the first, the BHK kidney cell line was transfected with Mi in the presence or absence of a 6× histidine–ubiquitin-encoding plasmid. Cellular lysates were passed over a nickel resin to bind His-tagged ubiquitin, and bound proteins were analyzed by Western blot using anti-Mi antibody. As shown in Figure 3A, Mi was identified in pull-downs only from lysates in which tagged ubiquitin and Mi had been expressed together. A limitation of this experiment was that misexpression of Mi in kidney cells may not reflect events in pigment cells.

Therefore, ubiquitination of endogenous Mi was tested with or without Sl stimulation of human melanoma cells transfected with HA-tagged ubiquitin. Cellular lysates were immunoprecipitated with Mi antibody, resolved by SDS-PAGE, transferred to nitrocellulose, and probed for ubiquitin with anti-HA antibody. A significant increase in Mi/ubiquitin conjugates was detected after stimulation by Sl [Fig. 3B]. In both of these experiments it is notable that some endogenous MAPK activity exists even in the absence of exogenous Steel factor addition, likely explaining the presence of modest but detectable ubiquitination of Mi in unstimulated MeWo.

Figure 2. c-Kit-induced Mi degradation is blocked by proteasome inhibition. Presence of a Mi dephosphorylating activity. Mi degradation kinetics were followed using cycloheximide (CHX). (A) Human melanoma cells [501 Mel] were stimulated by recombinant Steel factor [SI] in the presence of cycloheximide [SI + CHX] and with and without the proteasome inhibitor MG132. α-Tubulin was probed as internal loading control. (B) Lactacystin was used as proteasome inhibitor on SI-stimulated human melanoma cells. (C) Primary human melanocytes were stimulated with recombinant SI in the presence of cycloheximide [SI + CHX] in the absence or presence of MgI32. Reappearance of the lower migrating Mi isoform at later time points was reversed by treatment of cells with the phosphatase inhibitor okadaic acid in the human melanoma MeWo. (D) Mouse melanoma cells [B16] were treated with TPA to activate MAPK as these cells lack functional c-Kit receptor. Mi levels were compared in the absence or presence of MgI32 at the time points indicated. (E) Mouse melanocytes were treated with TPA as above and Mi levels assessed by Western blotting.
These studies show that Mi (either endogenous or exogenous) can be modified covalently by ubiquitin in response to SL/c-Kit signaling.

Ser-73 is dispensable for Mi degradation

The presence of functional Mi was next measured by assessing DNA-binding activity in nuclear extracts of melanoma cells after stimulation by SL, with or without proteasome inhibition. To assess simultaneously the role of specific phosphorylation and degradation of Mi, wild-type and specific Mi mutants were engineered into adenoviral expression vectors and used to infect (>90%) melanoma cells efficiently before examination of DNA-binding activity in nuclear extracts. Virally expressed, epitope-tagged Mi-specific DNA binding was identified by monoclonal antibody supershift in an electrophoretic mobility shift assay (EMSA) using radiolabeled DNA probe. Mi DNA-binding activity declined within 2 hr in cells exposed to SL, whereas no loss of DNA binding was seen in cells treated with SL plus MG132 (Fig. 4, left). Because c-Kit signaling results in Mi phosphorylation at ser-73, the role of this serine residue in destabilization was examined. As previously shown (Price et al. 1998a), DNA-binding activity is retained with mutation of ser-73 to alanine. However, the S73A and wild-type proteins were found to be equally labile after SL stimulation (Fig. 4, right). Furthermore, DNA-binding activity of the S73A mutant was also stabilized by proteasome inhibition. These results indicate that although MAPK phosphorylation of ser-73 occurs after SL stimulation, it is not absolutely required for SL/c-Kit-induced destabilization of Mi. Therefore, another consequence of c-Kit signaling must contribute to the observed Mi destabilization.

Because SL/kit-induced degradation of Mi is blocked

Figure 3. Mi is covalently ubiquitinated in a c-Kit responsive fashion. (A) Plasmids encoding Mi or His-tagged ubiquitin were transfected transiently into baby hamster kidney cells (BHK) and cell lysates were made 24 hr later. His-tagged protein (and associated protein) was affinity purified on nickel resin, proteins eluted and resolved by SDS-PAGE, and Western blotted for Mi. (B) HA-tagged ubiquitin plasmid was transfected into human melanoma cells (as indicated). Twenty-four hours later cells were stimulated with recombinant Steel factor (SL, as indicated) and extracted 0.5 hr later. Lysates were immunoprecipitated with anti-Mi antibody followed by Western blotting for HA (ubiquitin). Endogenous Mi covalently associates with ubiquitin in a SI-inducible fashion.

Figure 4. Ser-73 is not essential to c-Kit-induced Mi degradation. Adenoviruses encoding wild-type Mi or ser-73 mutated to alanine (S73A) were used to infect human melanoma cells (501 Mel), which were subsequently stimulated with Steel factor (SL) in the presence or absence of cycloheximide and MG132 proteasome inhibitor. Nuclear extracts from these cells were used in gel shift assays and the Mi-specific DNA-binding activity was identified by supershifting with anti-HA antibody addition (as indicated). Both wild-type and S73A mutant Mi DNA-binding activities diminish after SL stimulation.
by MAPK inhibition (Fig. 1), the mediators of this signal must reside at the level of MAPK or downstream. One possible explanation for the unperturbed degradation of the S73A mutant was that a second phosphorylation could contribute to ubiquitin-dependent proteolysis. Mi contains a number of potential phosphorylation targets but no additional MAPK consensus. One of these, S409, resides within a consensus sequence for the Rsk family of serine/threonine kinases (RRXS/T) [Leighton et al. 1995]. Rsk also represents a known downstream target of MAPK activation [Tsai et al. 1993; Dalby et al. 1998; Gavin and Nebreda 1999], potentially explaining why MEK inhibition could stabilize Mi in the context of Sl treatment.

**Phosphorylation of Mi by Rsk-1 in response to c-Kit stimulation**

To test whether Rsk may phosphorylate Mi directly at ser-409, initial studies examined in vitro kinase (IVK) activity. As shown in Figure 5A, immunoprecipitates of Rsk-1 from Sl-stimulated melanoma cells could phosphorylate recombinant Gst–Mi fusion protein, but not a Mi fusion carrying the S409A mutation. This activity was strongly induced by Sl stimulation of the melanoma cells. Evidence of autophosphorylation by Rsk in response to c-Kit stimulation was also seen, as indicated (Fig. 5A). An unidentified substrate protein that coprecipitates with Rsk is indicated by an asterisk. The kinetics of Rsk and MAPK/ERK activation after Sl stimulation were also examined in side-by-side IVK reactions followed by PhosphorImager quantitation. Peak Rsk activation paralleled but slightly lagged the induction of MAPK/ERK activity in melanoma cells after Sl stimulation (Fig. 5B), as anticipated for a kinase that is, itself, a substrate for (and activated by) MAPK [Blenis 1993]. Therefore, Rsk-1 is capable of directly phosphorylating recombinant Mi on ser-409 in vitro in a manner regulated by Sl stimulation with kinetics that recall those of the ser-73 Mi kinase ERK.

To determine whether Rsk-1 and Mi interact in vivo, we asked whether the endogenous forms of the two pro-

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**Figure 5.** Involvement of Rsk-1 in phosphorylation of ser-409. (A) In vitro kinase assays were carried out using immunoprecipitated Rsk-1 [or protein A control] from unstimulated or Steel factor-stimulated [Si] melanoma cells. Kinase substrates consisted of GST [control], GST fusion to carboxy-terminal fragment of wild-type Mi ["Mi"], or GST fusion to carboxy-terminal fragment of Mi mutated at ser-409 to alanine ["S409A"]. GST blot is shown at the bottom. Rsk-1 autophosphorylation is indicated and a weakly coprecipitating second Rsk substrate is indicated by asterisk. (B) Kinetics of Rsk and MAPK/ERK activation after Si stimulation. Kinase assays as described in A were quantitated by PhosphorImager and plotted as percent of maximal activity, indicating that Rsk activation lags ERK activation slightly. (C) Rsk-1 associates with Mi upon Si stimulation. Human melanoma cells were stimulated with Sl [or unstimulated] in the presence or absence of MEK inhibitor drug [PD98059]. Cell lysates were either probed directly for p90–Rsk-1 or Mi [lanes 1–3]. MEK inhibition prevents a Steel-induced mobility shift in both Mi and a subtle mobility shift in Rsk-1. Mi was also immunoprecipitated from cell lysates followed by Western blotting for Rsk-1. Rsk-1 associated with Mi after Steel stimulation only if MAP kinase was not inhibited. (D) Two-dimensional phosphotryptic mapping indicates presence of phosphoserine 409 within endogenous cellular Mi. The two-dimensional mobility of phosphoserine 409 was determined by analyzing in vitro phosphorylated recombinant Mi as in Fig. 5A. 32P-labeled melanoma cells were stimulated by Sl and extracted; endogenous Mi was immunoprecipitated and subjected to tryptic digestion and two-dimensional analysis (simultaneously run with in vitro phosphorylated recombinant Mi). A spot was observed [dotted circles], which displays superimposable migration relative to the phosphoserine 409 spot generated by Rsk-1 phosphorylation of ser-409 in vitro.
tein. Lysates of SI-stimulated [presence or absence of MEK inhibitor] or unstimulated human melanoma cells were immunoprecipitated with Mi antibody and examined by Western blot for coprecipitation of Rsk. As shown in Figure 5C, the mobility shift of Mi is prevented by MAPK inhibition [lanes 1–3]. Probing of the same blot for Rsk-1 reveals equal abundance of the protein, although with some evidence for a slight mobility shift with SI stimulation [Fig. 5C, lanes 1–3]. From these same lysates, endogenous Rsk-1 specifically comminoprecipitates with endogenous Mi only after SI stimulation [lanes 4,5]. Moreover, MAPK inhibition prevents this Rsk-1:Mi association [lane 6]. Thus, endogenous Mi and Rsk proteins appear to physically associate with one another after SI stimulation in a manner that is dependent on MAPK activation.

We then examined whether endogenous Mi was phosphorylated at ser-409 in vivo using two-dimensional phospho-tryptic mapping. To identify the ser-409 phosphopeptide, recombinant Mi was phosphorylated in vitro by Rsk-1 as above, and a tryptic digest was resolved by electrophoresis and thin layer chromatography using an apparatus producing inverted replicas (Fig. 5D). For comparison, endogenous cellular 32P-labeled Mi protein was immunoprecipitated from melanoma cells, digested, and resolved in the same way. Several 32P-labeled spots were apparent, one of which perfectly comigrates with the ser-409 Rsk-phosphorylated recombinant species based on superimposing the images [encircled in Fig. 5D]. Although these experiments cannot directly prove that Rsk phosphorylates Mi in vivo, several observations support this possibility: [1] the presence of a Rsk target consensus site at ser-409, [2] the stimulation of endogenous Rsk activity in response to SI with the same kinetics as those of ERK; [3] the physical association of Rsk with endogenous Mi in response to SI stimulation; [4] the profound stabilization of Mi in response to MAPK inhibition; and [5] the presence of a phosphopeptide from endogenous Mi that is indistinguishable from a Rsk-phosphorylated Mi peptide. These observations are consistent with the possibility that Rsk phosphorylates Mi in cells at ser-409 in response to SI signals.

Double mutations of MAP kinase and Rsk target serines profoundly stabilize Mi

Mutation of S409 to alanine was examined for its impact on the signal-dependent proteolysis described above. NIH 3T3 cells were transfected with wild-type Mi, single serine-to-alanine mutants at positions 73 or 409 [S73A and S409A], or a double mutant at both serines 73 and 409 [S73/409A], and stabilities were tested with stimulation with TPA. As shown in Figure 6A, wild-type protein and the two single point mutants were all degraded rapidly. In contrast the double mutant at both S73 and S409 was significantly stabilized. Similarly, transfection into human melanoma cells followed by c-Kit stimulation produced short-lived wild-type and single mutant proteins [S73A or S409A], whereas the S73/409A double mutant was profoundly stabilized [Fig. 6B].

Coupled activation/degradation: S73/409A mutant Mi is transcriptionally incompetent although intact for DNA binding

Transactivation potentials of the mutant Mi proteins were assessed by transfection/reporter assay. This is of particular importance because SI stimulation has been shown previously to superactivate Mi function, and mutation of ser-73 partially abrogates transactivation [Hemesath et al. 1998]. The current observations suggest that the identical modification that activates Mi also targets the protein for degradation. If so, the stabilized double mutant S73/409A might be a poor transcriptional activator despite its greater stability. To test this, Mi expression plasmids were cotransfected into NIH 3T3 as well as human melanoma cells with a luciferase reporter gene driven by the tyrosinase promoter, a Mi-responsive target [Bentley et al. 1994; Hemesath et al. 1994; Yasumoto et al. 1994; Bertolotto et al. 1996]. As shown in Figure 7A, wild-type Mi stimulated the tyrosinase promoter, in agreement with prior results [Hemesath et al. 1998]. The S73A and S409A single mutants were less efficient transactivators than wild-type Mi. However, the double mutant was devoid of transactivation potential despite its greater stability. The lack of activity for the double mutant is specific for this reporter, as all data
are normalized to a cotransfected constitutive reporter. Of note, the double mutant reproducibly diminishes promoter activity below the basal (vector control) levels, consistent with the possibility that it acts dominant negatively with respect to other E box factors in these cells. If the double mutant exhibits dominant negative function, it would be expected to retain DNA-binding activity, as predicted from the location of the two mutations (which are remote from the bHLHZip motif).

DNA-binding activity from nuclear extracts of cells transfected transiently with either wild-type Mi or the S73/409A double mutant were examined by EMSA after MAPK stimulation. The Mi/DNA complex was again identified using supershifting monoclonal antibody. Whereas wild-type Mi displayed a significant loss of DNA-binding activity after MAPK stimulation, the S73/409A double mutant was resistant to this reduction in DNA binding (Fig. 7B). DNA-binding patterns closely recapitulated the direct protein levels seen by Western blot after transfection of the Mi constructs. These data indicate that mutations of serines 73 and 409 do not perturb DNA binding of the Mi protein, and that the stability seen with the double mutant in protein blots is reflected in functional protein isolated from the nuclei of cells. Because simultaneous mutations of both serines 73 and 409 were necessary to produce Mi protein resistant to this degradation pathway, phosphorylation at either ser-73 or ser-409 in response to c-Kit stimulation appears to be sufficient to signal ubiquitin-dependent proteolysis of Mi. These experiments demonstrate that simultaneous mutation of serines 73 and 409 prevents signal-induced Mi degradation and produces a stable protein, but at the same time cripples the transactivation potential of Mi. Thus, the signals that produce transcriptional activation and protein degradation of Mi are functionally coupled in melanocytes.

Discussion

The data presented here suggest that the Sl/c-Kit signaling pathway in melanocytes targets the transcription factor Mi simultaneously for activation and proteolytic degradation. This conclusion is based on evidence that the half-life of Mi is shortened by c-Kit-triggered MAPK activity. Because MAPK inhibition profoundly stabilizes Mi protein, the necessary events that signal Mi degradation appear to reside at or downstream of MAPK. On the basis of mutational and functional studies these include MAPK itself and likely Rsk-1 (although other kinases might potentially substitute for Rsk-1; see below). These kinases appear to phosphorylate Mi at serines 73 and 409, respectively, either of which is sufficient to target Mi for degradation. Sl stimulation induces the association of Rsk-1 with Mi, and Rsk-1 activity is measurably up-regulated in these cells with a kinetic profile that closely follows that of MAPK. Covalent modification by ubiquitin also suggests that the degradation pathway involves ubiquitin-dependent proteolysis, likely mediated by proteasome activity that was repressible by several proteasome inhibitors. Because simultaneous alanine substitutions at both serines 73 and 409 produce highly stabilized, but transcriptionally inactive Mi, the stimulation and degradation signals appear to be biochemically coupled.

The existence of such tightly coupled activation/degradation signals likely creates a circumstance in which up-regulatory stimuli are received at target gene promoters, while simultaneously limiting their duration. This
potential scenario is of interest, in part, because of the multiple activities with which Mi is associated. For example, in contrast to the net degradation of Mi after Sl/c-Kit stimulation, the MSH signaling pathway profoundly up-regulates Mi gene expression, producing significantly elevated Mi protein levels over many hours (Bertolotto et al. 1998a; Price et al. 1998b). Whereas Mi protein may be depleted with Sl/c-Kit stimulation, prolonged transcription of the Mi gene after MSH results in sustained levels, despite the presence of constitutive basal MAPK activity (Price et al. 1998b). Thus, the short-lived effects of Sl/c-Kit signaling on Mi may provide a mechanistic basis for analyzing the diverse roles of Mi within melanocytes.

One implication of Mi degradation after Sl/c-Kit stimulation is that in tumor cells (melanomas) pathologic c-Kit activation could produce net diminished Mi levels and function. This prediction is of interest because of numerous reports that c-Kit expression is frequently absent in melanomas, and that Steel factor stimulation of melanomas, which retain c-Kit, may result in paradoxically antiproliferative effects (Zakut et al. 1993). However, intermittent subcutaneous Steel factor injection into humans clearly induces melanocytic proliferation (Costa et al. 1996), as predicted from the known function of c-Kit as a mitogenic receptor tyrosine kinase and proto-oncogene in other contexts (Williams et al. 1990).

The net inhibition of melanoma growth upon sustained c-Kit stimulation is a behavior that may correlate with Mi degradation. A recent clinical series (King et al. 1999) demonstrated nuclear Mi expression in 100% of 76 consecutively accessioned melanoma specimens, despite concomitant loss of other melanocytic markers such as pigment, c-Kit, pmel/17 (HMB 45), and other melanocytic markers (Carrel and Rimoldi 1993; Zakut et al. 1993; Halaban et al. 1997; Kaufmann et al. 1998). Because Mi plays an essential post-developmental role in melanocyte proliferation or survival (based on the Mi"mutant), it is possible that degradation of Mi with persistent c-Kit stimulation might be inhibitory to melanoma growth.

The observation that activation and degradation appear to be coupled to the identical phosphorylation events on Mi raises the question of how phospho-Mi is recognized for degradation. The above data demonstrate a role for ubiquitin-mediated proteolysis and proteasomal activity. It is interesting to speculate that p300/CPB, which is selectively recruited to Mi by MAPK-mediated phosphorylation (Price et al. 1998a), might in some manner directly facilitate Mi degradation as well. Such a role for p300/CPB has been suggested in the regulation of p53 degradation (Grossman et al. 1998). This could involve p300/CPB-dependent recruitment of a separate ubiquitin ligase or might reflect an intrinsic ubiquitin ligase activity of p300/CPB itself. It has been suggested that signal mediated degradation of a transcription factor bound to a promoter is required for dissociation of the pro-initiation complex and transcription elongation (Nawaz et al. 1999). Mechanistically, it is possible that promoter-bound, phosphorylated Mi may eventually impede transcriptional initiation or even elongation, and for this reason it could be advantageous to degrade [or dephosphorylate] the activated protein. Such a model could operate for many signal-responsive transcriptional pathways.

Another paradoxical observation, which might be explained by the coupled activation/degradation signals described here, involves regulation of pigmentation. Mi recognizes and, in experimental systems, has been shown to transactivate the tyrosinase, TRP-1, and TRP-2 pigment enzyme gene promoters (Bentley et al. 1994; Hemesath et al. 1994; Yasumoto et al. 1994; Bertolotto et al. 1996, 1998b). This transactivation activity is measurably enhanced by MAPK-mediated activation of Mi (Hemesath et al. 1998). However, dominant active Ras has been found to diminish pigmentation and dominant negative Ras to increase it (Englaro et al. 1998) despite evidence that c-Kit up-regulates Mi through MAPK (Hemesath et al. 1998) and can enhance pigmentation in vivo (Costa et al. 1996). One explanation for these findings is that the same signaling pathway may target Mi for activation as well as proteolytic degradation. Homocystine down-regulation is a hallmark of receptor tyrosine kinase signaling and occurs at multiple levels including cytokine/receptor internalization, dephosphorylation, and proteolysis. The current study extends such homocystine down-regulation to a nuclear target of c-Kit.

Mi phosphorylation at ser-73 is strongly established as a target of MAPK/ERK on the basis of in vitro and in vivo biochemical and inhibitor data (Hemesath et al. 1998). Significant correlative data suggest that Rsk-1 may target ser-409, although lack of Rsk inhibitor reagents renders this point incompletely proven in vivo. Moreover, ser-409 might serve as a target substrate for other kinases, such as protein kinase A (PKA), whose consensus sequence overlaps. Indeed, in vitro kinase assays demonstrate the ability of PKA to phosphorylate ser-409 (data not shown). Although phosphorylations of serines 73 and 409 may be coupled in the c-Kit signaling pathway, it is possible that phosphorylation at ser-409 may occur independently of MAPK activation in other signaling contexts, and such events should be sufficient to destabilize the Mi protein.

The kinases ERK-2 and p90 Rsk are physically associated with one another in the cytoplasm of many cell types (Scimeca et al. 1992) including 501Mel cells (data not shown) and translocate to the nucleus together upon MEK-mediated activation of ERK. ERK has been shown to phosphorylate Rsk and stimulate the activity of its amino-terminal kinase domain (Sturgill et al. 1988; Bennis 1993; Grove et al. 1993). We found that the kinetics of ERK and Rsk activation in response to c-Kit signaling closely paralleled one another in melanoma cells. This is the first example of these two regulated kinases acting in concert to modulate the stability and the cofactor coupling of a sequence-specific transcription factor. The only other common phosphorylation target of the kinases ERK and Rsk is c-Fos, which together with c-Jun forms the transcription factor complex AP-1. Phosphorylations at the extreme carboxyl terminus of c-Fos by ERK and Rsk may result in an enhancement of the growth-
promoting properties of c-Fos in fibroblast cells [Chen et al. 1993, 1996], although the exact molecular mechanisms are unclear.

IκB, a cytosolic inhibitor of the transcription factor NFκB, undergoes phosphorylation-dependent ubiquitin-mediated degradation in response to a number of extracellular stimuli. Rsk is one of several kinases that may be involved in triggering NFκB nuclear translocation by the phosphorylation of ser-32 in IκB, targeting the latter for degradation [Ghoda et al. 1997, Schouten et al. 1997]. There are other kinases triggered by distinct receptor systems that serve the same function in NFκB induction. Ubiquitin-mediated degradation of Stat proteins also produce signal-responsive regulation for this transcription factor family [Kim and Maniatis 1996].

The transcriptional effects observed with phosphorylation of Mi in this system appear to involve p300/CREB coactivator recruitment [Sato et al. 1997; Price et al. 1998a]. It is important to consider, however, that the involvement of these coactivators may vary at different transcriptional targets. Thus, some promoters may use Mi for p300-dependent transcription, whereas others might not require p300 for Mi-driven transcription. The discovery of transcriptional targets genes of Mi, particularly in the contexts of these distinct signaling pathways, may shed important light on both the biological output of the signals and the mechanisms whereby Mi as a common transcriptional intermediate may modulate diverse functional pathways in human disease.

Materials and methods

Cell culture and treatments

Human melanoma cell lines 501 MEL and MeWo (gifts of Dr. Ruth Halaban, Yale Medical School, New Haven, CT; Zakut et al. 1993) were grown in F10 medium [GIBCO-BRL] with 10% fetal bovine serum plus penicillin/streptomycin/glutamine [GIBCO-BRL]. Human primary melanocytes [gift of Dr. Ruth Halaban, Yale Medical School, New Haven, CT; Zakut et al. 1993] were cultured in MBM2 medium according to the manufacturer’s protocol. Murine melanocytes (gift of Dr. Ruth Halaban, Yale Medical School, New Haven, CT; Zakut et al. 1993, 1996), although the exact molecular mechanisms are unclear.

Microphthalmia signaling and degradation

501 MEL cells were also infected with E1A- and E1B-deleted adenovirus type 5, which contained human wild-type MITF and ser-73 to alanine-mutated MITF, respectively, both of which were HA tagged. The MITF insert was cloned into a unique XbaI site located 452 bp upstream from the 5’ end of the vector backbone. The insert was driven by an elongation factor α promoter [Mizushima and Nagata 1990]. Subconfluent 501 MEL cells were incubated with the adenovirus constructs in serum-free F10 supplemented with 10 mM MgCl₂ for 30 min at a multiplicity of infection of 100. After the infection the medium was replaced by fresh F10 medium plus 10% fetal bovine serum, and cells were cultured for another 62 hr followed by human Steel factor stimulation in the presence or absence of cycloheximide and the proteasome inhibitor MG132.

Immunoblotting and immunoprecipitation

The monoclonal antibody D5 was raised against a histidine fusion protein expressed from the amino-terminal Taq–Sac fragment of human Mi cDNA [Tachibana et al. 1994] and produces a specific supershift of a Mi/DNA complex, but does not react with any closely related family members (not shown). An alternative Mi monoclonal antibody C5 [Weilbaecher et al. 1998] was used to detect murine Mi. High-affinity rat HA antibody [Boehringer Mannheim] was used to detect HA-tagged Mi protein, and antibody against α-tubulin (Sigma) for protein-loading control. For immunoblot analysis, cells were lysed in lysis buffer [50 mM Tris [pH 7.6], 150 mM NaCl, 1% Triton-X 100] plus protease inhibitors [Boehringer Mannheim] and phosphatase inhibitors [20 mM NaPP, 10 mM NaF, and 1 mM Na₃VO₄].

Protein concentrations were determined by using the Dc protein assay kit [Bio-Rad]. Samples were solubilized in SDS sample buffer plus 50 mM DTT and boiled for 5 min. After SDS-PAGE and transfer to nitrocellulose, blots were blocked in 5% milk plus 0.05% Tween-20 in Tris-buffered saline before antibody incubation. Antibody complexes were detected with peroxidase-conjugated secondary antibody [Cappel] and chemiluminescence reagents [Amersham].

For immunoprecipitation, cells were lysed with lysis buffer as above or with RIPA buffer [10 mM Tris-HCl [pH 7.5], 150 mM NaCl, 1% Nonidet P-40, 0.1% SDS, 1% sodium sarcosyl, 1 mM DTT] plus protease inhibitors. The soluble fraction was incubated overnight at 4°C with primary antibodies and subsequently protein G agarose beads [GIBCO-BRL] were added to the solution and incubated for an additional 1 hr at 4°C. Beads were washed three times with cold PBS or RIPA buffer, resuspended in SDS sample buffer, and boiled for 5 min. The eluted proteins were resolved on SDS-PAGE and immunoblot analysis performed as described above. For His-tagged protein analysis, washed Ni-NTA agarose beads [Qiagen] were added to the cell lysates and incubated at 4°C for 4 hr. Eluted proteins were analyzed as above.

Supershift EMSA

Cells were washed twice in cold PBS and resuspended in buffer A [10 mM HEPES [pH 7.9], 1.5 mM MgCl₂, 10 mM KCl, and 0.5 mM DTT plus protease inhibitors] and incubated on ice for 10 min. The incubations were agitated for 10 sec and centrifuged
(14,000 rpm for 10 sec using a microcentrifuge), and the resulting pellet was resuspended in two packed-nuclei volumes of buffer C [20 mM HEPES (pH 7.9), 420 mM NaCl, 1.5 mM MgCl2, 0.2 mM EDTA, 0.5 mM DTT, 25% glycerol] plus protease inhibitors and further incubated on ice for 30 min. The suspension was centrifuged and the supernatant was quantitated, and equal amount of protein used to perform gel mobility assay. Probe labeling and DNA-binding reactions were performed as described [Hemesath et al. 1998]. For supershift EMSA, 1 µl of high affinity HA antibody [Boehringer Mannheim] was added concurrently with other components of the binding reaction.

Transfection assay

Transfections of 501 MEL or 3T3 cells were carried out in 24-well plates using FuGENE 6 as above according to manufacturer’s recommendations. Each experiment was performed in triplicate, and transfection efficiency was normalized to cotransfected sea pansy luciferase plasmid (Promega). Cells in each well were transfected with 0.1 µg of tyrosinase reporter [Hemesath et al. 1998], 0.5 µg of MITF expression plasmid or its derivative mutant vectors, and 0.1 µg of sea pansy luciferase plasmid. After 48 hr of transfection, cells were lysed in 150 µl of passive lysis buffer as per manufacturer’s recommendations [Promega] and incubated for 30 min at room temperature. An aliquot of the lysates was used to perform luciferase assays using dual luciferase reporter system (Promega). Cells in each well were transfected with 0.5 µg of MITF, 0.5 µg of MITF expression plasmid or its derivative mutant vectors, and 0.1 µg of sea pansy luciferase plasmid. After 48 hr of transfection, cells were lysed in 150 µl of passive lysis buffer as per manufacturer’s recommendations [Promega] and incubated for 30 min at room temperature. An aliquot of the lysates was used to perform luciferase assays using dual luciferase reagents [Promega]. An equal amount of lysate was subjected to immunoblotting analysis for expression of the MITF (wild type or its derivatives).

Phosphotryptic mapping

For in vivo labeling of MITF, 501 MEL cells were washed in phosphate-free, serum-free DMEM (GIBCO) and then labeled in vivo with 1 mCi/ml of inorganic 32P in a 10-cm dish for 8 hr. Cells were washed with ice-cold PBS, then lysed in 1% Triton, 170 mM NaCl, 20 mM Tris (pH 7.4) plus protease and phosphatase inhibitors. Lysates were immunoprecipitated with anti-MITF monoclonal antibody [CS] [Weilbaecher et al. 1998] and protein G agarose beads (GIBCO) at 4°C overnight. Beads were washed and boiled in loading buffer (3% SDS, 10% glycerol, 120 mM Tris, and 0.1 M DTT). Proteins were separated on SDS-PAGE and transferred to nitrocellulose membranes. Bands were visualized by autoradiography, excised, and digested for 20 hr at 37°C with 25 µg of TPCK-treated trypsin (Sigma). Phosphopeptide mapping was carried out as described [Hemesath et al. 1998] with the Pharmacia LKB Multiphor II apparatus, which generates inverted replica blots, and using ammonium carbonate (pH 8.9).

For two-dimensional phosphotryptic analysis of in vitro Rsk-1 labeled MITF, a recombinant fragment of MITF fusion protein spanning amino acid residues 370–419 was used. 501 MEL cells were starved in serum-free medium for 2 hr and then stimulated with 20 ng/ml of recombinant human Steel factor (R & D Systems). Cells were washed with ice-cold PBS, then lysed in 1% Triton, 170 mM NaCl, 20 mM Tris (pH 7.4) plus protease and phosphatase inhibitors. Lysates were immunoprecipitated with polyclonal antibody to Rsk-1 (Santa Cruz) and protein G agarose beads (GIBCO) at 4°C overnight. Beads were washed twice in PBS and 10 µCi of [32P]ATP were added to 100 ng of recombinant GST–MITF in 20 µl of IVK buffer [50 mM HEPES (pH 7.6), 10 mM MgCl2, 2 mM NaVO4, 0.5 mg/ml Pefabloc, 2 mM DTT, and 50 µM ATP] and incubated at 32°C for 30 min. Loading buffer was added to the sample, boiled, then resolved on SDS-PAGE, transferred to nitrocellulose, and subjected to trypptic digestion and two-dimensional analysis as described above.

In vitro kinase assay

Cells were Steel-stimulated and lysed as described above and 4 µl of anti-Erk-2 or anti-Rsk-1 antisera (Santa Cruz) plus protein G agarose beads were added to the lysates and mixed overnight at 4°C. Beads were washed three times with lysis buffer and once with IVK buffer [50 mM HEPES (pH 7.6), 2 mM NaVO4, 10 mM MgCl2, 1 mM PMSF, 2 mM DTT, and 50 µM ATP]. For each reaction, 440 µl of IVK buffer, 10 µCi [32P]ATP, and recombinant phosphoacceptor proteins were added. Mi histidine fusion proteins spanning residues 16–185 containing ser-73 (Tachibana et al. 1994) or GST fusion protein spanning residues 389–419 containing ser-409 were added as substrates and incubated at 30°C for 30 min. Reactions were stopped by addition of loading buffer and analyzed by immunoblotting and autoradiography.

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