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Ligand-dependent Degradation of SRC-1 is Pivotal for Progesterone Receptor Transcriptional Activity

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**Abbreviations** : NR, Nuclear Receptor; PR, Progesterone Receptor; SRC-1, Steroid Receptor Coactivator-1; bHLH, basic Helix-Loop-Helix; LB, leptomycin B; NES, Nuclear export signal; PRE, Progesterone Response Element; SERM: Selective Estrogen Receptor Modulator.
ABSTRACT

The progesterone receptor (PR), a ligand-activated transcription factor, recruits the primary coactivator SRC-1/NCoA-1 to target gene promoters. It is known that PR transcriptional activity is paradoxically coupled to its ligand-dependent down-regulation. However, despite its importance in PR function, the regulation of SRC-1 expression level during hormonal exposure is poorly understood.

Here we report that SRC-1 expression level (but not other p160 family members) is down-regulated by the agonist ligand R5020 in a PR-dependent manner. In contrast, the antagonist RU486 fails to induce down-regulation of the coactivator and impairs PR agonist-dependent degradation of SRC-1. We show that SRC-1 proteolysis is a proteasome- and ubiquitin-mediated process that, predominantly but not exclusively, occurs in the cytoplasmic compartment where SRC-1 colocalizes with proteasome antigens as demonstrated by confocal imaging. Moreover, SRC-1 was stabilized in the presence of leptomycin B or several proteasomal inhibitors. Two degradation motifs, amino-acids 2-16 corresponding to a PEST motif and amino-acids 41-136 located in the bHLH domain of the coactivator, were identified and shown to control the stability as well as the hormone-dependent down-regulation of the coactivator. SRC-1 degradation is of physiological importance since the two non-degradable mutants that still interacted with PR as demonstrated by co-immunoprecipitation, failed to stimulate transcription of exogenous and endogenous target genes, suggesting that concomitant PR/SRC-1 ligand-dependent degradation is a necessary step for PR transactivation activity. Collectively, our findings are consistent with the emerging role of proteasome-mediated proteolysis in the gene regulating process and indicate that the ligand-dependent down-regulation of SRC-1 is critical for PR transcriptional activity.
INTRODUCTION

The progesterone receptor (PR), also known as NR3C3, plays a crucial role in the coordination of several aspects of female reproductive development and function (1). Invalidation of the PR gene in mice leads to pleiotropic reproductive abnormalities and demonstrates that PR orchestrates key events associated with the establishment and maintenance of pregnancy. From a pathophysiological perspective, accumulating evidence indicates that PR is involved in breast cancer cells proliferation and is implicated in the development and progression of breast cancer (2). Coregulators (coactivators or corepressors) are important Nuclear Receptor (NR)-recruited cofactors modulating NR-mediated transcription leading to activation or repression of target specific genes (3). SRC-1 is a PR coactivator belonging to the p160 gene family which contains three homologous members (SRC-1, -2, and -3) serving as NR transcriptional coactivators (4). This family of coactivators is characterized by the presence of several conserved functional domains: a bHLH-PAS N-terminal domain, a CBP interacting domain (AD1), a glutamine-rich region, a C-terminal activation domain (AD2), and several LXXLL boxes involved in NR binding. The p160 coactivators are defined as “primary coactivators” whose activity is regulated by posttranslational modifications (5-10). The current models indicate that p160 coactivators serve as a recruitment platform for other coactivator complexes carrying intrinsic enzymatic activities to specific enhancers/promoters leading to the covalent modification of specific histones and/or other coregulators involved in the transcriptional machinery (11, 12).

Several experiments have revealed a tight association between the turnover rate of several NR and their transcriptional activity, showing that both aspects of NR function appear to be inversely related (13-18). Among the factors regulating PR levels are its ligands. It was initially shown that administration of progesterone to ovariectomized guinea pigs provoked a rapid fall in uterine receptor concentration (19). Hormone-dependent down-regulation of PR has been finally confirmed by several groups (20-22) but its biological significance is still unclear. Phosphorylation of PR on a key serine residue (Ser294) by MAPKs was shown to couple multiple receptor functions, including ligand-dependent PR down-regulation by the ubiquitin-proteasome pathway (13). The concept that transcriptional activation and ubiquitin-mediated proteolysis are interdependent processes is emerging as a potentially important control mechanism of transcription (16, 23). Although their significance
remains to be defined, it appears that complex interactions between regulatory molecules governing both transcription and ubiquitination/degradation exist (24-26). However, little is known concerning the fate of coregulators during ligand-dependent NR down-regulation (27, 28).

In a previous study, we have shown that SRC-1 is exported from the nucleus to the cytoplasm and speculated that this export might be a regulatory mechanism controlling the termination of hormone action possibly through its degradation (29). In order to establish a link between SRC-1 proteolysis and the PR-mediated transcription process, we studied the mechanism governing SRC-1 proteolysis at the steady-state level and questioned whether the ligand could modulate its turn-over. In this study, we demonstrate that SRC-1 undergoes covalent modifications by ubiquitin which targets the coactivator to the proteasome at the steady-state level. We identify two critical degron domains directly linked to the coactivator proteolysis. Aside from this ligand-independent stability regulation, we show that SRC-1 undergoes accelerated agonist-dependent and PR-mediated down-regulation via the ubiquitin–proteasome pathway. SRC-1 proteolysis occurs concomitantly of ligand-dependent PR degradation. Of note, the nature of the ligand is shown to be critical for this process since both PR and SRC-1 ligand-dependent proteolysis was inhibited in the presence of RU486, leading to dramatic loss of PR transactivating capability.
RESULTS

_SRC-1 mainly colocalizes with cytoplasmic proteasome antigens_

In our previous report about the regulatory mechanisms of SRC-1 subcellular trafficking, we have shown that SRC-1 localizes both in nuclear and cytoplasmic corpuscular structures (29). Several studies have reported coregulators localization in organelles (30, 31). We tried to identify the nature of these cytoplasmic and nuclear speckles by colocalization studies with various antigens and with fluorescent organelles markers. Since several nuclear receptors (NR) and coactivators such as SRC-3 have been shown to interact with the proteasome (32, 33), we used confocal microscopy to investigate whether proteasome components might also accumulate in SRC-1 speckles. By using antibodies directed against the human S7 subunit of the 19S (Rpt1) and the α/β subunits of the 20S proteasome, we found that SRC-1 colocalized with both 26S proteasome antigens (Fig 1A and Supplemental Fig S1). The fluorescence intensity profile indicates that colocalization was predominant in SRC-1 speckles: simultaneous fluorescence intensity increase was observed in cytoplasmic speckles but also in lesser extent in nuclear speckles (Fig 1B), suggesting that SRC-1 is mainly but not exclusively proteolyzed in the cytoplasm. Similar intensity profiles were obtained for cells immunolabeled for SRC-1 and the 20S proteasome (data not shown). A partial colocalization of SRC-1 was also found with the Promyelocytic Leukemia Protein (PML) in the typical nuclear domain (ND10) (Supplemental Fig S2 A). Such an association has been already described (34). In contrast, nuclear speckles did not overlap with transcription sites as evidenced by the absence of colocalization with the SC-35/SRp30 spliceosome component (Supplemental Fig S2 B). Similarly, no colocalization of SRC-1 with organelles like mitochondria, lysosomes, peroxisomes or the Golgi apparatus could be observed (Supplemental Fig S2 C-E and data not shown).

_SRC-1 is ubiquitinylated in vivo and is degraded by the proteasome_

We next studied the mechanism of SRC-1 down-regulation. First we investigated whether the coactivator was ubiquitinylated and targeted to the proteasome. COS-7 cells were transfected with the expression vector encoding the full-length SRC-1 and incubated in the presence of proteasome inhibitors, MG132 or epoxomicin. Consistent with previous reports (14, 35), both inhibitors increased
SRC-1 protein level in comparison to cells treated with vehicle (Fig 2A and Supplemental Fig S3). To demonstrate that SRC-1 is poly-ubiquitinylated, COS-7 cells were transfected with SRC-1 expression vector in the presence or absence of a vector encoding His-tagged ubiquitin (His 6-Ub) and analyzed by Western Blot. In the absence of His 6-Ub, the anti-SRC-1 antibody detected a major band of ~160 kDa (Fig 2B, left panel, lane 1). In cells cotransfected with His 6-Ub expression vector, a moderate decrease in band intensity was observed with a slightly higher molecular weight smear, indicative of ubiquitinylated moieties (Fig 2B, left panel, lane 2). His-tagged proteins were purified by chromatography on nickel-charged agarose beads (Ni-NTA) and analyzed by Western Blot with an anti-SRC-1 antibody to show that these bands correspond to ubiquitinylated SRC-1 (Fig 2B, right panel, lane 2).

Several cytoplasmic proteasome substrates have been shown to relocalize into the nucleus upon stabilization by proteasome inhibitors (36-38). We thus examined whether SRC-1 subcellular distribution was similarly modified in such conditions. Indeed, overnight treatment of cells with MG132 induces an obvious shift of the coactivator into the nucleus (Fig 2C). This result suggests that escape from cytoplasmic proteolytic degradation stimulates the nuclear accumulation of SRC-1 (36).

If our hypothesis is true, then inhibition of SRC-1 nuclear export should induce SRC-1 expression level stabilization. To verify this point, we followed the turnover rate of SRC-1∆(NES), a mutant deleted of its nuclear export signal (NES) (29). The result shows a better stability of this mutant compared to the wt SRC-1 (Supplemental Fig. S4). In a similar approach, we used the nuclear export inhibitor leptomycin B (LB) to impede wt SRC-1 access to cytoplasm. In presence of LB, SRC-1 not only relocalized into the nucleus [data not shown and (29)] but its expression level also increased ~2.5 fold (Fig 2D). However, SRC-1 stabilization with LB did not reach the level obtained with MG132 (data not shown; and compare quantification Fig 2A to Fig 2D). Thus, the nuclear accumulation of the coactivator indicates a possibility of a partial degradation of SRC-1 in the nuclear compartment. Interestingly, similar experiments with the p160 coactivator SRC-3, which has been shown to be degraded mainly in the nucleus (39) showed no significant increase of SRC-3 expression level under LB treatment (Fig 2D). Overall, our data show that SRC-1 turnover is a proteasome- and ubiquitin-mediated process that takes place, predominantly but not exclusively, in the cytoplasm.
Agonist ligand enhances concomitant proteolysis of PR and SRC-1

We next studied SRC-1 degradation in the context of PR activation. Progestins are known to induce PR proteolysis by the proteasome (22, 40). In addition, Li et al have shown that upon ligand treatment, progesterone receptor (PR) preferentially interacts with SRC-1 (41). We thus investigated whether SRC-1 down-regulation might be also modulated by PR ligands. As previously reported (22), immunocytochemical studies (Fig 3A) and Western Blot experiments (Supplemental Fig S5) showed that the agonist ligand R5020 stimulates stably expressed endogenous PR proteolysis after 24 h treatment while the antagonist ligand RU486 prevents PR proteolysis in Ishikawa cells stably expressing PR-B (Ishi-PR-B). To test the impact of ligands on SRC-1 expression level, Ishi-PR-B cells were transiently transfected with a SRC-1 expression vector and incubated overnight with R5020 or RU486. Western Blot analyses revealed that SRC-1 and PR are concomitantly degraded in the presence of agonist R5020 and that RU486 prevents the degradation of both proteins (Fig 3B). Similar results were obtained using different Ishi-PR-B subclones (data not shown). Real time quantitative RT-PCR excluded the possibility of any ligand-dependent down-regulation of SRC-1 mRNA levels (Supplemental Fig S6). MG132 exposure inhibited the agonist-dependent proteolysis of SRC-1 (Fig 3B, lane 4), indicating that this stimulated down-regulation is mediated by the proteasome. Importantly, using antibodies specifically detecting endogenous SRC-1, we similarly observed agonist-dependent degradation of endogenous SRC-1 in Ishi-PR-B cells (Fig 3C and Fig 3D). Of note, a 10-fold excess of antiprogestin RU486 abrogated the R5020-dependent degradation of endogenous SRC-1 and PR as shown in Fig 3D (third lane), suggesting that SRC-1 degradation is tightly linked to the ligand-dependent PR activation. To further verify this hypothesis, we tested if SRC-1 proteolysis could be stimulated in the absence of PR. We used the Ishikawa parental cell line (Ishi-PR-0) initially used to establish the Ishi-PR-B cell line and that lacks PR-B expression (42). Ishi-PR-0 cells were transfected with SRC-1 expression vector and incubated 24 h with R5020 or RU486. Under these conditions, both ligands did not affect SRC-1 expression level, indicating that SRC-1 down-regulation requires the presence of PR-B (Fig 3E). Finally, we determined whether other p160 coactivators such as SRC-2/TIF2/GRIP-1 or SRC-3/AIB1, which are also known proteasome targets (14), could be degraded in response to R5020. None of these coactivators was significantly degraded under similar
experimental conditions (Supplemental Fig S7), suggesting a target-specific coactivator effect of PR.

It has been initially proposed that antiprogestins are capable of inducing PR down-regulation but with much slower kinetics than agonists (22). We therefore tried a longer time point to check if SRC-1 degradation was occurring in presence of RU486. The result shows that, in contrast to 24 h incubation (Fig 3F, lane 3), 48 h treatment with RU486 induced a significant reduction of both SRC-1 and PR (Fig 3F, lane 4). More importantly, in presence MG132, RU486 treatment resulted in a dramatic accumulation of PR and SRC-1 (Fig 3F, lane 5), showing that RU486-induced down regulation is mediated by the proteasome. Thus, these results not only indicate that RU486 impairs the ligand-dependent down-regulation of PR and SRC-1 by slowing down their degradation, but also confirm the concomitance of their ligand-dependent proteolysis. Collectively, our results indicate that specific SRC-1 turn-over is modulated in a ligand-dependent manner and requires PR expression.

Identification of SRC-1 domains involved in its degradation

In order to elucidate the mechanisms driving SRC-1 to the proteasome under basal conditions, we identified the domains involved in SRC-1 turn-over. In silico analysis of SRC-1 primary sequence was carried out in search for putative PEST degradation motifs. The result indicated that amino-acids 2 to 16 of SRC-1 had a high score (+9.63) for this type of motif. We therefore focused our investigation on the N-terminal subdomain of the coactivator. A critical importance of the bHLH domain for AIB1/SRC-3 mediated proteolysis has been previously reported by Li and colleagues (39) . Thus, we also explored the role of this domain in SRC-1 down-regulation. Two deletion mutants were generated lacking either the PEST sequence, or the bHLH domain, encompassing amino-acids 2 to 16 [Δ(PEST)] and amino-acids 41 to 136 [Δ(bHLH)], respectively (Fig 4A).

In order to investigate if these two motifs were involved in SRC-1 degradation, wt SRC-1, Δ(PEST) or Δ(bHLH) mutants were expressed in COS-7 cells, and cycloheximide was added to block protein neosynthesis. The decay of wt SRC-1 and mutant proteins was monitored and quantified by western blot as a function of time. SRC-1 expression levels decreased after 1 h and almost disappeared after 6 h (Fig 4B, left panel), indicating of a half-life of approximately 3 h. In contrast, both Δ(PEST) and
∆(bHLH) expression levels showed no decrease under the same experimental conditions (Fig 4B, middle and right panel), showing that ∆(PEST) and ∆(bHLH) mutants are more stable than wt SRC-1. To confirm that these motifs were involved in proteasome-mediated SRC-1 degradation, we compared both mutants and wt SRC-1 localization by immunocytochemistry and found that in contrast to the wild-type coactivator (Fig 1) and the ∆(PEST) mutant, the ∆(bHLH) mutant localized predominantly in the nucleus (Fig 4C). In contrast to the wild-type coactivator (Fig 1), colocalization studies of both mutants with 19S proteasome antigens S7/Rpt1 and with the α/β proteasome 20S subunits showed no significant overlap (Fig 4C and data not shown).

Moreover, to investigate the involvement of these domains on SRC-1 protein stability, we compared the impact of MG132 on both mutants with wt SRC-1. While SRC-1 protein levels were increased ~3 fold under 15 h MG132 treatment (Fig 4D, left panel), the expression level of either ∆(PEST) or ∆(bHLH) remained unchanged under the same conditions (Fig 4D, middle and right panels). Similarly, expression levels of both mutants were not increased in presence of epoxomicin (Supplemental Fig S8). Of note, quantification comparison of band intensity (Fig 4D, histograms) showed that both mutants were expressed to a greater extent than the wild-type coactivator, suggesting that the deletions may have indeed a stabilizing effect on these mutants. Taken together, our observations show that amino-acids 2-16 and 41-136 are involved in SRC-1 down-regulation by targeting SRC-1 to proteasome degradation at the steady-state.

N-terminal degradation motifs of SRC-1 are necessary for its ligand-dependent down-regulation

In order to evaluate the contribution of the two degradation domains in the context of the hormonal activation, we transiently transfected Ishi-PR-B cells with either wt SRC-1, ∆(PEST) or ∆(bHLH) mutants. We hypothesized that if the two degradation motifs are also involved in hormone-stimulated down-regulation of SRC-1, then both mutants should not undergo proteolysis under hormone stimulation. As expected, after 24 h of R5020 treatment, wt SRC-1 was significantly down-regulated, while the expression level of both mutants showed no significant variation (Fig 5A). Interestingly, the ligand-dependent down-regulation of PR still occurred in each condition, showing that the receptor
down-regulation does not require SRC-1 degradation (Fig 5A). To exclude the possibility that the two deletions may have impaired the interaction between the SRC-1 and PR, we conducted reciprocal co-immunoprecipitation experiments in cells transiently expressing PR and either wt SRC-1 or the deletion mutants. The result shown in Fig 5B indicates that PR reciprocally co-immunoprecipitates with wt SRC-1 as well as with Δ(PEST) and Δ(bHLH) mutants. Taken together, these results indicate that under hormonal stimulation, SRC-1 ligand-dependent proteolysis requires both degradation signals.

Since we showed that SRC-1 could be partially proteolyzed in the cytoplasm where it colocalized in speckles with the proteasome (Fig 1), we next wondered if PR will colocalize in the same cytoplasmic speckles. This may specially be the case if we consider the work of Qiu et al. who have shown that PR down-regulation under hormone treatment occurs in the cytoplasm (43). Our result shows that in the absence of hormone, SRC-1 is expectedly cyto-nuclear and does not colocalize with PR (Fig 6A). Eight hours of hormonal treatment (in the presence of cycloheximide) induces the nuclear accumulation of both PR and SRC-1, indicative of their interaction during the nuclear import (29). Interestingly, the ligand also induces the colocalization of PR and its coactivator in cytoplasmic speckles (Fig 6A), suggesting that PR/SRC-1 complexes might be exported back to the cytoplasm. In contrast, in the presence of R5020, Δ(PEST) and Δ(bHLH) mutants were efficiently accumulated in the nucleus, consistent with our coimmunoprecipitation data showing that they do interact with PR in the presence of ligand, but did not colocalize with PR in cytoplasmic speckles (Fig 6B-D). Overall, this experiment suggests that PR and SRC-1 could be proteolyzed as a PR/SRC-1 complex through the same proteasome.

Ligand-dependent proteolysis of SRC-1 is necessary for PR-mediated gene transactivation

To examine the functional link between SRC-1 degradation and its coactivating function, we investigated the impact of coactivator proteolysis on PR-mediated transcription. To this aim, we first analyzed if the proteasome function was required for efficient PR transcriptional activation. Cotransfection of PRE2-TATA-luc reporter gene with the PR encoding vector was performed in parental Ishi-PR-0 cells (devoid of PR), either alone or in combination with the vector encoding SRC-
Twenty-four hours after transfection, cells were treated for 24 h with R5020 alone or in combination with MG132. To exclude the possibility that the cellular toxicity of MG132 might affect general transcription in Ishikawa cells, we used a 500 nM concentration of the inhibitor, a dose compatible with cell survival of endometrial carcinoma cell lines (44). We show that MG132 drastically attenuates ligand-dependent PR transactivation (Fig 7A), confirming previous observations made by Dennis et al (45). Interestingly, SRC-1-potentiated PR-mediated transcription was also abolished by the proteasome inhibitor (Fig 7A). This result suggests that the proteasome-mediated degradation is required not only for PR transcriptional activity but also for SRC-1-potentiation of PR. To further explore the relationship between coactivator degradation and the functional consequences on PR-mediated transcription, we used the two non degradable mutants Δ(PEST) and Δ(bHLH) in cotransfection experiments with PR (Fig 7B). Since these 2 mutants are not efficiently degraded by the proteasome (see Fig 4B and 4D), we predicted that they might not exert efficient potentiation of PR transactivation. Indeed, in the presence of R5020, SRC-1 strongly coactivated PR while both Δ(PEST) and Δ(bHLH) mutants were unable to enhance PR-mediated transactivation as compared to wt SRC-1 (Fig 7B). These results suggest that the concomitant degradation of SRC-1 and PR is necessary for efficient transcriptional activity of the receptor. Finally, to determine whether the functional link between SRC-1 proteolysis and its coactivating properties were also relevant for human endogenous gene activation, we quantified the level of the progesterone-induced amphiregulin gene that we have previously studied (46). Parental Ishi-PR-0 cells were transfected with PR alone or in combination with wt or SRC-1 mutants. Amphiregulin mRNA levels were significantly increased upon R5020-dependent PR activation and were further enhanced in the presence of SRC-1 (Fig 7C). Conversely, coexpression of PR with either Δ(PEST) or Δ(bHLH) mutant significantly reduced amphiregulin expression (P <0.001). Taken together, our results demonstrate that hormone-induced degradation of SRC-1 is physiologically relevant for potentiation of PR-mediated transcriptional events.
DISCUSSION

In this study, we investigated the impact of SRC-1 proteolysis on PR-mediated transcription. We provided evidence that the agonist-dependent degradation of SRC-1 is pivotal for PR-mediated transcription. We have established that agonist ligand R5020, but not antagonist RU486, induces the concomitant degradation of endogenous or ectopic PR and SRC-1. Interestingly, SRC-1 turnover requires the presence of PR. Both basal and induced SRC-1 down-regulation are mediated through the proteasome pathway and seem to occur at least in part, in the cytoplasmic compartment. Two regions located in the N-terminal part of SRC-1 (i.e., a PEST motif and amino-acids 41-136 of the bHLH domain) were identified as two degron motifs. Both signals were shown to be responsible for basal- and hormone induced-degradation of SRC-1. Deletion of each of these domains [Δ(PEST) and Δ(bHLH) mutants] leads to non-degradable SRC-1 mutants insensitive to proteasome inhibitors. By comparing the biological functions of these two mutants with wt SRC-1, we found that they were incapable of potentiating PR-mediated transactivation on a synthetic PR response-element but also on amphiregulin, an endogenous PR target gene. The HAT motif and the CBP interacting domain of SRC-1 are known to regulate the transcriptional activity of SRC-1 (47, 48). Both regions are present in Δ(PEST) and Δ(bHLH) mutants (Fig 4A), and therefore the reduced PR-dependent transactivation of the mutants is not due to an alteration of these regulatory domains but rather to a defect in down-regulation. Thus, our results are indicative of a functional link between proteasome-mediated down-regulation of SRC-1 and its coactivating property.

We have previously shown that SRC-1 is a transcriptional coactivator whose localization is hormonally regulated in the presence of PR (29). Mainly functioning in the nuclear compartment, this coactivator may also be present in the cytoplasm, predominantly concentrated in cytoplasmic speckles (29). Several studies have also demonstrated that p160 coregulators might be localized in the cytoplasm (7, 30, 31). Although the concentration of SRC-1 in cytoplasmic speckles was initially reported to be linked to overexpression (49), it has been also observed for endogenous p160 coactivators (50) and, more importantly, a recent study correlated this archetypical distribution with the cytoplasmic sequestration of SRC-1 by SIP (SRC-Interacting Protein) (51). During our primary
search to identify the nature of these speckles, we initially observed a colocalization between SRC-1 and proteasome antigens, indicating that SRC-1 cytoplasmic speckles are enriched of proteasome components (Fig 1A). Similar subcellular distribution studies already reported SRC-2 colocalization with proteasome antigens but specifically at the nuclear level (34, 52). Coactivator/proteasome interaction have been also described at the biochemical level for the p160 coactivators (33, 53), as well as NR such as the thyroid receptor, the retinoic acid receptors RARα and RXR, the estrogen receptor ERα, or the vitamin D receptor (32). We detected a strong colocalization in the cytoplasmic compartment although a weaker colocalization in speckles was also observed in the nuclear compartment (Fig 1B) indicative of a predominant but not exclusive proteolysis of the coactivator in the cytoplasmic compartment. Interestingly, nuclear export of SRC-3 has been shown to be required for its proteasomal degradation (54). However, our finding is not consistent with the work of Li et al. who recently showed that proteasome-dependent turnover of SRC-3 occurs specifically in the nucleus (39). Although we could not completely exclude that nuclear degradation also occurs for SRC-1 (see colocalization profiles Fig 1B), this discrepancy between SRC-1 and SRC-3 argues for the fact that each SRC family member has different and specific physiological functions (55).

We have shown that the ubiquitin-proteasome pathway mediates selective degradation of SRC-1 and regulates the steady-state expression level of the coactivator. Similarly, Yan et al have shown that several SR coactivators were degraded through the ubiquitin-proteasome dependent pathway and that SRC-1 proteolysis occurs specifically through the Ubiquitin-Conjugating Enzyme 2 (35). The half-life regulation of p160 coactivators has been extensively investigated since the discovery of their prototype SRC-1 and several studies have demonstrated the physiological and pathophysiological importance of regulating SRC-1 expression levels (56-58). SRC-1 is an important modulator of PR-mediated gene transcription and in order to accurately exert its physiological function its level must be therefore tightly regulated in vivo. In this context, Han et al used an original transgenic mouse model in which SRC-1 levels were shown to influence the compartment specific corepressor-to-coactivator ratio in order to modulate PR activity in uterus (59). Cell regulation of SRC-1 levels seems to be also critical for tumorigenesis and studies have demonstrated that SRC-1 expression is significantly increased in breast tumors and positively correlates with disease recurrence and poor disease-free
survival (55). Consistent with this finding, SRC-1 level is up-regulated during mammary tumor progression (60) and the role of this coactivator in promoting mammary tumor cell invasion was recently demonstrated in vivo (57, 58).

Beside the regulation of SRC-1 proteolysis at basal level, the present study also analyzes ligand-stimulated down-regulation of the coactivator. Similarly to other rapidly turned over transcription factors, engagement of PR in transactivation has been shown to be coupled to PR degradation by the ubiquitin-proteasome pathway (13). However, the functional impact of the SRC-1 coactivator on PR-mediated transactivation has never been clearly established. We demonstrate for the first time that concomitantly to PR degradation, SRC-1 proteolysis is dramatically increased in the presence of the agonist ligand R5020 and that this process is mediated through the proteasome. Similarly to PR (22), this down-regulation is necessary for PR-mediated transcription. Recent advances in molecular biology have redefined the role of proteasome as a regulatory system that influences the fate of many cellular processes, such as cell proliferation, apoptosis, and more recently gene transcription. Despite the disparate nature of the later process, a growing body of evidences indicates that ubiquitin and the proteasome are intimately involved in NR-mediated gene control (45, 61, 62). Steroid hormone receptors and their coactivators cycle onto and off steroid-responsive promoters in a ligand-dependent manner and it is now believed that the ubiquitin–proteasome functions in promoting the turnover of transcription complexes, thereby facilitating proper gene transcription (16, 63). Dennis et al. have proposed the existence of a transcriptional mechanism that link the proteasome function with the continued recruitment of RNAPII to sustain the transcriptional response (45). Consistent with these observations are the fact that (i) a number of ubiquitin pathway enzymes and components of the proteasome have been found to act as modulators of NR function (24, 26, 64) and that (ii) enzymes and components of the proteasome are recruited to the promoters of NR-responsive genes (16, 63).

In spite of this, it is difficult to conceive how a coactivator will be paradoxically part of a coactivating complex positively modulating gene activation and at the same time a specific target of the ubiquitin-proteasome pathway. Thus, the coupling of PR/SRC-1 proteolysis and efficient transcriptional activation is counterintuitive and rather puzzling but could be a general phenomenon occurring during transcription (65). Consistent with this, is the fact that neither PR nor its coactivator were down-
regulated in presence of the antagonist RU486. This result may suggest that RU486 indirectly prevents
recruitment of the proteasome machinery, thereby inhibiting transcription. The same observation was
made with ERα and the partial antagonist Tamoxifen though it may not be considered as a general
phenomenon for Steroid Receptor since the pure antagonist Faslodex dramatically stabilize ER in
similar conditions (49). It is not the first example of a hormonal regulatory mechanism implicated in
specific coregulators proteolysis: indeed, SRC-2 is down-regulated through the activation of the
cAMP dependent protein kinase pathway (52). More importantly, Gianni et al showed that SRC-3, but
not SRC-1 or SRC-2, is phosphorylated by p38MAPK in a Retinoic Acid-dependent manner and then
degraded by the proteasome pathway (27). In this case, phosphorylation of SRC-3 has a biphasic
effect on RARα transactivation with facilitation followed by restriction of transcription.
Since the presence of PR is required for SRC-1 degradation, two important remaining questions
concern the identification of the key-player responsible for SRC-1 degradation and whether this factor
is involved in both basal and ligand-induced SRC-1 down-regulation. Shao et al used RNA
interference to knock-down SRC-3 that consequently abolishes ERα ligand-dependent degradation,
suggesting that the coactivator itself regulates ERα degradation (66). Conversely, since the two non-
degradable mutants did not impede the ligand- induce PR down-regulation (Fig 5A), our results do not
converge towards a link between the recruitment of a common E3-ligase by SRC-1 which will in turn
induce the ligand-dependent degradation of the PR/SRC-1 complex. The signal that targets PR and
SRC-1 to progress from transcription to degradation may also involve post-translational modifications
operating like a molecular signature such as phosphorylation, ubiquitinylation or sumoylation (9, 67,
68). Alternatively, direct recruitment of ligase in the vicinity of the coactivator complex or directly at
the enhancer level may be also implicated in SRC-1 turnover along with PR. A good candidate would
be the PR-B coactivator/ubiquitin ligase E6-AP since this coactivator plays a major role in controlling
the regulated degradation of SRC-3 and PR-B isoform (54, 69). Alternatively, the colocalization with
proteasome antigens observed in our study (Fig 1) might also be linked to the direct interaction
observed between SRC-1 and the proteasome through the Low Molecular mass Polypeptide 2
proteasome subunit (LMP2) (53). Such a direct ligand-dependent interaction may drive the coactivator
to proteolysis. Another potential candidate for PR and SRC-1 degradation might be Jab1, a coactivator
involved in ER degradation (70). We are currently investigating this hypothesis, since we have shown in a previous study that Jab1 is a coactivator of PR, inducing the formation of a PR/SRC-1/Jab1 ternary complex during the transcription process (71).

In summary, we demonstrate in the present study that SRC-1 expression level is hormonally regulated by the ligand. While, in presence of an agonist the PR/SRC-1 complex is proteolyzed in order to achieve transcription, an antagonist as RU486 impairs the ligand-dependent degradation of PR/SRC-1 and consequently the transactivation process. Our data indicate that the expression level of SRC-1 coactivator is critical for PR transcriptional activity. These findings are consistent with the emerging role of the 26S proteasome in the gene regulation process (72). P160 family members are certainly not the only coactivators implicated in such processes and it will be interesting to elucidate the sequential progression of each coregulator degradation during gene regulation.
MATERIALS AND METHODS

Hormone and inhibitors

Cycloheximide, Epoxomicin (Epoxo), MG132, Leptomycin B (LB) were purchased from Sigma (St Louis, MO). Agonist R5020 (17,21-dimethyl-19-norpregna-4,9-dien-3,20-dione) and antagonist RU486 (Sigma, St Louis, MO) were used at a concentration of 10nM, except where indicated.

Plasmids

Nomenclature: derivatives denoted with a Δ lack the protein segment delineated by the numbered amino-acids. Plasmids encoding the wild-type human progesterone receptor (pSG5-PR) and coactivator SRC-1 (pSG5-SRC-1, pSG5-HA-SRC-1, pSG5-HA-GFP-SRC-1) have previously been described (29). PCR-based site-directed mutagenesis of pSG5-HA-SRC-1 was used to create deletion mutants: pSG5-HA-SRC-1-Δ(2-16) [named “Δ(PEST)"], pSG5-SRC-1-Δ(41-136) [named “Δ(bHLH)”] and pSG5-HA-SRC-1Δ(990–1060) [named “Δ(NES)”, (29)]. The plasmid pPRE2-TATA-Luc has been previously described (71). Plasmid pSG5-His6-Ub is a gift of D. Bohmann (Laboratory EMBL, Heidelberg). Plasmids pSG5-SRC-2 and pCR3.1-SRC-3 have been described previously described (7, 73) and GFP-Peroxisome targeting signal expression vector was purchased (Clontech, Mountain View, CA).

Cell culture and DNA transfection

Human endometrial Ishikawa cells (parental cell line “Ishi-PR-0” and stable “Ishi-PR-B”) were provided by Dr LJ. Blok (Erasmus University, Rotterdam, Netherland) (74). COS-7, HEK293, Ishi-PR-0 and Ishi-PR-B were grown in DMEM containing 10% fetal bovine serum (Biowest, Miami, FL) and supplemented with L-glutamine and antibiotics (penicillin / streptomycin, PAA Laboratories GmbH, Austria). For hormonal regulation experiments, cells were grown in the presence of 10% steroid-depleted FBS prior (24h) and during transfection experiments. Transfections were performed with the indicated expression vectors using LipofectAMINE 2000 according to the manufacturer’s recommendations (Invitrogen, Carlsbad, CA).
**Antibodies**

Monoclonal anti-PR antibodies used in the study were the Let126 (0.5µg/mL) (75), the monoclonal anti-PR from Novocastra (NCL-L-PGR-312/2) or the rabbit polyclonal anti-PR (sc-538) from Santa Cruz Biotechnology, used for immunoprecipitaion. Anti-SRC-1 mouse monoclonal antibody (Millipore, Billerica, MA) was used for Western Blot and immunocytochemistry (1µg/mL). Endogenous SRC-1 was detected with anti-SRC-1 (sc-6096) purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Anti-HA 3F10 (200ng/mL) was from Roche Applied Science (Indianapolis, IN). Rabbit polyclonal antibody directed against human S7/Rpt1 and 20S proteasome subunits, and KAT13C/NCOA2/SRC-2 were purchased from Abcam (Cambridge, MA) and used at 1:1000 dilution. Anti-α-tubulin (1:10000) and anti SC-35 (1µg/mL) were purchased from Sigma (St Louis, MO). Anti-PML was provided by H de Thé (IUH, Paris, France). Anti-SRC3/AIB1 antibody was purchased from BD Biosciences (San Diego, CA) and was used at 0.5µg/ml. Secondary antibodies (1:4000) : anti-mouse, anti-rat, anti-rabbit antibodies conjugated to alexa 488 (green) or 595 (red) or Dylight 549 (red) were from Invitrogen (Carlsbad, CA) and Jackson ImmunoResearch Laboratories (West Grove, PA). Secondary peroxydase-conjugated anti-mouse (Calbiochem, San Diego, CA) and anti-rabbit (Vector laboratories Inc., Burlingame, CA) antibodies were used at 1:15000 dilution.

**Luciferase reporter gene assays**

COS-7 cells were cultured in free steroid medium and reverse transfected in 96-well plates with 4ng PR, 100ng PRE2-TATA-Luciferase, 100ng SRC-1 (wild-type or mutants), and 5ng β-galactosidase (internal control). The pBlue-Script plasmid was used to equally adjust DNA quantity. After 24h transfection, cells were incubated with or without 10nM R5020 for 24h. Cells were collected with the Passive Lysis Buffer (Promega, Madison, WI) and luciferase activity was measured with a luminometer (Victor, Perkin Elmer, Waltham, MA). Luciferase activity was normalized with β-galactosidase activity. The results are means ± S.E. of four independent experiments.
Immunocytochemistry

Cells were seeded in 24-well plates and processed as previously described (7). Briefly, cells were fixed with 4% paraformaldehyde and permeabilized for 30 min with a 0.5% solution of Triton X100 diluted in PBS. Cells were then incubated with primary antibody overnight at 4°C, followed by the appropriate fluorochrome-coupled secondary antibody (alexa 488 or 595, Invitrogen; or Dylight 549, Jackson ImmunoResearch Laboratories) for 30 min. Nuclear counterstaining was performed with 0.5 µg/mL DAPI (4',6'-diamidino-2-phenylindole) and coverslips were mounted on slides with ProLong Gold mounting medium (Invitrogen, Carlsbad, CA). For standard microscopy (Fig 2 and 3), fluorescent cells were observed with an Olympus Provis AX70 and images were acquired with Qcapture Pro version 5.1 (Q Imaging Inc., Surrey, BC) using an Evolution VF Monochrome camera (Media Cybernetics Inc., Bethesda, MD).

Confocal Microscopy

For Fig 1, 4 and 6, a Zeiss LSM-510 confocal laser scanning microscope (Carl Zeiss, Thornwood, NY) was used for fluorescence acquisition. Images of fixed cells were collected from equatorial planes of cells with a pinhole setting of ~1.0 airy unit (AU) (optical thickness of 0.8 µm) using a x63:1.4NA oil immersion plan-apochromat objective with X8 frame averaging accumulation. In order to exclude crosstalk artifacts, both red and green fluorescence emission were acquired sequentially in separated channels. The confocal microscope settings were kept the same for all scans. To validate colocalization of proteins (Fig 1 and 4), line scans of intensity profiles across the cells were generated with the LSM browser software (76). This function associates the merge images with an intensity profile of each channel, measured along a freely positioned line. To obtain an average representative intensity profile expressed as arbitrary units (AU), lines were drawn through the middle of each cell images in a distance covering the cytosol and the nucleus. Green lines represent the intensity profile for the proteasome antigen S7/Rpt1 signal and the red lines represent the intensity profile for SRC-1 signal.
**Western Blot and Immunoprecipitation**

Cells were lysed in lysis buffer (50 mM Tris-HCl [pH 8.0], 150 mM NaCl, 1% Triton X-100, 2 mM EDTA, 0.2 mM NaF, 0.2 mM Na₃VO₄, protease inhibitor cocktail) for 15 min, and the debris were cleared by centrifugation at 14000 × g for 15 min at 4°C. Samples were resolved by 7.5% SDS gel electrophoresis and transferred onto nitrocellulose membranes. The indicated antibodies were diluted in TBST buffer supplemented with 5% non-fat milk and added to the membranes for 1h30 at room temperature (RT) or overnight at 4°C followed by incubation with the appropriate horseradish peroxidase-conjugated secondary antibodies for 45 min at RT. All proteins were detected with ECL Plus detection reagents (Amersham Biosciences Corp, Piscataway, NJ) and visualized by chemiluminescence. For the normalization, the membrane was stripped, probed with anti-α-tubulin antibody diluted to 1:1000 (Sigma, St Louis, MO). The bands were quantified after digitalization on a gel scanner using Image J software. Results, mean of 3 independent experiments (except Fig 5A), are presented as the ratio SRC-1(or PR)/α-Tubulin and are expressed as fold induction above the value measured for wild-type SRC1 in the absence of MG132 arbitrary set at 1. For coimmunoprecipitation, HEK 293 cells were transfected in 100mm plate with either wt SRC-1, Δ(PEST) or Δ(bHLH) plasmids, and cultured in presence of 10⁻⁸M R5020 for 24h. Cells were lysed at 4°C in 500µl lysis buffer and cell debris were pelleted by centrifugation (14.000 rpm, 15 min, 4°C). Immunoprecipitation of the supernatant with anti-SRC-1 or with the rabbit polyclonal anti-PR or with IgG control were performed with Protein G Magnetics Beads (Millipore, Billerica, MA) according to the manufacturer instructions. Bound immunocomplexes were boiled in Laemmli buffer, separated by 7.5% SDS-PAGE, blotted nitrocellulose membranes with anti 1µg/mL SRC-1 (Millipore, Billerica, MA) and anti PR-B (Let 126, 0.5µg/mL) antibodies, detected with ECL Plus detection reagents (Amersham, Biosciences Corp, Piscataway, NJ), and visualized by chemiluminescence.

**Real Time RT-PCR**

The Ishikawa cell line expressing PR-B or not was transfected by the indicated plasmids by Polyfect reagent (QIAGEN, Valencia, CA) in six-well plates (six wells per condition). After a 2 h-treatment by R5020 10nM, cells were washed and lysed by Trizol reagent (Life Technologies, Gaithersburg, MD).
Total RNA were extracted as described by the manufacturer. One microgram of each sample was treated by DNase I and was reverse transcribed using random primers as previously described (77). Real-time quantitative PCR of amphiregulin gene was performed as described (46) using the Power SYBR Green master mix (Applied Biosystem, Carlsbad, CA) in duplicate with 1:20 fraction of each cDNA sample and the corresponding primers, using an ABI Prism 7300 apparatus. For each sample, the mRNA concentration was extrapolated from standard curve and averaged Ct value was divided by that of the corresponding reverse-transcribed 18S RNA (relative mRNA).

Statistical analysis

Data are expressed as the mean ± SEM. Mann Whitney U-test was used to determine significant differences between two groups. For multiple comparisons, Kruskal-Wallis test followed by Dunn’s post-test was performed using the computer software Prism 4 (GraphPad Software, San Diego, CA). Statistical significance is indicated at P values < 0.05, 0.01 and 0.001.
**FIGURES LEGENDS**

**Fig.1. Colocalization of SRC-1 with the 26S proteasome by confocal microscopy.**

A, Colocalization analysis between HA-SRC-1 and endogenous proteasome antigens S7/Rpt1 and 20S subunits. COS-7 cells were transiently transfected with the expression vector encoding HA-SRC-1. Cells were fixed after 40 h, immunolabeled with anti-HA and either anti-S7/Rpt1 or anti-20S antibodies, and then observed by confocal microscopy.

B, Validation of colocalization by scan of intensity profiles of a representative cell (expressed as arbitrary units, AU). Fluorescence intensity was calculated and plotted by drawing a line through the middle of the cell image in a distance covering several cytosolic and nuclear foci. Green lines represent the intensity profile for the proteasome antigen S7/Rpt1 signal and the red lines represent the intensity profile for SRC-1 signal. Indicated numbers refer to identified speckles: cytoplasmic (1 to 9), nuclear (8 to 11). Note that although the fluorescence intensity from the two channels is different, the peaks of both signals are overlapping.

**Fig.2. SRC-1 is proteolyzed by the 26S proteasome in a ubiquitin-dependent manner.**

A, COS-7 cells were transfected with the expression vector encoding SRC-1 and incubated in the absence or presence of MG132 (5 µM) during 15 h. Expression of SRC-1 was analyzed by Western blot using anti-SRC-1 and anti-α-tubulin antibodies. Bands intensity corresponding to SRC-1 were quantified as described in “Materials and Methods”.

B, CV-1 cells were transfected with the expression vector encoding HA-SRC-1 in the presence of the His6-tagged ubiquitin expression vector (His 6-Ub). Whole cell extracts were analyzed by electrophoresis on 6.4% SDS-PAGE and immunoblotted with anti-HA monoclonal antibody. Alternatively, the same co-transfected CV-1 cells were lysed in buffer containing guanidium-HCl (Ni-NTA). The ubiquitin-modified proteins were purified using Ni-NTA agarose beads as described under “Materials and Methods.” Affinity purified proteins were separated by electrophoresis, and His6-SRC-1 conjugates were detected by Western blot using the anti-HA monoclonal antibody. The ubiquitin conjugates of SRC-1 are indicated with brackets.

C, COS-7 cells were transfected with the expression vector encoding HA-SRC-1. Twenty hours post-
transfection, cells were incubated during 24 h with MG132 (1 µM) or treated with vehicle (DMSO). Cells were then fixed and immunolabeled with anti-HA antibody.

D, COS-7 cells were transfected with the expression vector encoding SRC-1 or SRC-3 and treated similarly than in C except that MG132 was replaced by Leptomycin B treatment (LB, 20 ng/ml). Expression levels of SRC-1 and SRC-3 were analyzed by Western blot using anti-SRC-1 or anti-SRC-3 monoclonal antibodies as indicated. Bands intensity representing the mean of at least 2 independent experiments were quantified as described in “Material and Methods”.

**Fig.3. Ligand- and PR-dependent SRC-1 proteolysis.**

A, Ishi PR-B cells, a cell line stably expressing PR-B, were cultured 24 h in the absence or in the presence of either the agonist R5020 (10 nM) or the antagonist RU486 (10 nM). Cells were then treated for immunocytochemistry with anti-PR antibody (Let 126) and observed by fluorescence microscopy.

B, Ishi PR-B cells were transfected with the SRC-1 encoding vector. After 48 h, cells were cultured 15 h as indicated, either in the absence of ligand (control vehicle, -H), in the presence of R5020 (10 nM), RU486 (10 nM), or in the presence of both R5020 (10 nM) and MG132 (5 µM). Whole cell extracts were analyzed by electrophoresis on 7.5% SDS-PAGE and immunoblotted with the indicated antibodies.

C, Non-transfected Ishi PR-B cells were treated as in B. Cells were immunolabeled for endogenous SRC-1 using an anti-SRC-1 antibody. Note the agonist-ligand-dependent down-regulation of endogenous SRC-1.

D, Non-transfected Ishi PR-B cells were cultured 24 h in the absence of ligand (vehicle, -H) or in the presence of either the agonist R5020 (10 nM) alone or in combination with a 100x excess of the antagonist RU486 (1 µM). Whole cell extracts were analyzed by electrophoresis on 7.5% SDS-PAGE and immunoblotted to detect endogenous SRC-1 and PR with the indicated antibodies.

E, Ishi PR-0 cells (parental cell line, devoid of PR) were treated as in Fig 3A. Cells were then analyzed by electrophoresis on 7.5% SDS-PAGE and immunoblotted with the indicated antibodies.

F, Ishi PR-B cells were transfected with the SRC-1 encoding vector. After 24h, cells were cultured
either in the absence of ligand (vehicle, -H), treated with R5020 (10 nM, 24h), RU486 (10 nM, 24 h
or 48 h), or RU486 (10 nM, 24 h) along with MG132 (1 µM). Whole cell extracts were analyzed by
electrophoresis on 7.5% SDS-PAGE and immunoblotted with the indicated antibodies.

**Fig.4.** The N-terminal Region of SRC-1 targets the coactivator to degradation.

A, Schematic representation of the wild-type coactivator SRC-1 (1441 amino-acids in length) with
boxes corresponding to major functional domains: bHLH: basic Helix Loop Helix domain, PAS: Per-
ARNT-Sim motif, NR1 and NR2: Nuclear Receptor-Interacting Domains 1 and 2, CBP/p300
interacting domain, Q: glutamine-rich domain. SRC-1 deletion mutants Δ(PEST) and Δ(bHLH) are
represented below with a thick line interrupted by a gap corresponding to the deleted amino-acids.

B, COS-7 cells were transfected as indicated with SRC-1, Δ(PEST) or Δ(bHLH) encoding vectors.

Seventy-two hours after transfection, cells were treated with cycloheximide (100 µg/ml) during 1, 4 or
6 h. Whole cell extracts were analyzed by electrophoresis on 7.5% SDS-PAGE and immunoblotted
with the indicated antibodies. Bands intensities (right panel) representing the mean of at least 2
independent experiments were quantified as described in “Material and Methods”.

C, Upper panel: Colocalization analysis of SRC-1 deletion mutants and S7/Rpt1. COS-7 cells were
transiently transfected with Δ(PEST) or Δ(bHLH) encoding vectors. Cells were fixed after 40 h and
immunolabeled with anti-HA and anti-Rpt1/S7 antibodies prior analysis by confocal microscopy.

Lower panel : scan of intensity profiles expressed as arbitrary units, AU. Fluorescence intensity was
calculated and plotted by drawing a line through the middle of the cell image in a distance covering
several cytosolic and nuclear foci. Green lines represent the intensity profile for the proteasome
antigen S7/Rpt1 signal and the red lines represent the intensity profile for Δ(PEST) or Δ(bHLH)
signals. Note the absence of significant peaks with overlapping signals.

D, COS-7 cells were transfected with HA-SRC-1, Δ(PEST) or Δ(bHLH) encoding vectors. After 48 h,
cells were incubated during 15 h with MG132 (5 µM) or vehicle. Whole cell extracts were analyzed
by electrophoresis on 7.5% SDS-PAGE and immunoblotted with the indicated antibodies. The band
intensities (right panel) were quantified as described in “Materials and Methods”. 
Fig 5. Ligand-dependent down-regulation of SRC-1 requires both degradation motifs of the coactivator.

A, Ishi PR-B cells were transfected as indicated with HA-SRC-1, Δ(PEST) or Δ(bHLH) encoding vectors. After 48 h, cells were cultured in the absence of ligand (vehicle, -), or in the presence of the agonist R5020 (10 nM) during 24h. The corresponding whole cell extracts were analyzed by electrophoresis on 7.5% SDS-PAGE and immunoblotted with the indicated antibodies. The band intensities (lower panel) were quantified as described in “Materials and Methods”.

B, HEK293 cells were cotransfected with PR and either the SRC-1, Δ(PEST), or Δ(bHLH) encoding vectors. Twenty four hours after transfection, cells were treated during 24 h with the agonist R5020 (10 nM). A coimmunoprecipitation assay was performed using either the anti-SRC-1, the anti-PR, or the IgG1 control antibodies (IgG1). Purified proteins were separated on 7.5% SDS-PAGE. Co-precipitated complexes were identified with the indicated antibodies.

Fig. 6. Colocalization of PR and SRC-1 in cytoplasmic speckles.

A, COS-7 cells were transiently transfected with the expression vector encoding HA-SRC-1 and PR. Twenty-four hours after transfection, cells were incubated or not for 8 h with R5020 in presence of cycloheximide (100 µg/ml) prior fixation. Cells were immunolabeled with anti-PR (Let 126) and anti-HA antibodies.

B and C, cells were treated as in A, except that PR was transfected as indicated with Δ(PEST) and Δ(bHLH), respectively.

D, Quantification of cells treated as described in A, B and C. Percent of cells treated with R5020 showing nuclear localization with or without cytoplasmic speckles. At least 100 cells were counted.

Fig.7. SRC-1 degradation is necessary for PR transcriptional activity.

A, Ishi PR-0 cells were cotransfected as indicated with expression vectors encoding PR and SRC-1 together with the reporter gene PRE2-TATA-luc and the internal control pRS-β-gal. Cells were incubated with R5020 (10 nM) and treated or not with MG132 (500 nM) during 24 h. Luciferase
activity was quantified and normalized by β-galactosidase activity. Data represent means ± SEM of at least three independent determinations.

B, COS-7 cells were cotransfected as indicated with HA-SRC1, Δ(PEST) or Δ(bHLH) encoding vectors, together with expression vector encoding PR, the reporter gene PRE2-TATA-luc and the internal control pRS-β-gal. Cells where treated during 24 h with R5020 (10 nM) or vehicle (control, -). Luciferase activity was quantified and normalized by β-galactosidase activity. Data represent means ± SEM of four independent determinations performed in triplicate.

C, Ishi PR-0 cells were cotransfected as indicated with HA-SRC-1, Δ(PEST) or Δ(bHLH) encoding vectors together with PR encoding vector and were treated with the agonist R5020 10 nM for 3 h. Total RNAs were extracted and relative expression of amphiregulin gene was quantified by qRT-PCR. Results, normalized by the amplification of 18S RNA, are mean ± SEM of three independent determinations. Statistical significance *** P<0.001 vs wild-type SRC-1 used as reference.

ACKNOWLEDGMENTS

The authors are indebted to Youssef Alj for the construction of the Δ(PEST) and Δ(bHLH) SRC-1 mutants. We thank C Massaad and BW O’Malley for kindly providing SRC-2 and SRC-3 plasmids, respectively. We gratefully acknowledge Luc Outin for excellent technical support and Geri Meduri for critical and thorough reading of the manuscript. We are thankful to Meriem Messina for her help in plasmid preparation.


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Fig. 2

A

160 kDa  
SRC-1  
50 kDa  
α-Tubulin

MG132  -  +  
Relative Fold  1  3.8

B

Whole cell extract  Affinity purified extract

SRC-1  +  +  
His 6-Ub  +  +  +  +  +

SRC-1  +  +  
α-Tubulin  -  -

C

HA-SRC-1  DAPI  MERGE

MG132  -  +  +  +

D

SRC-1  SRC-3

α-Tubulin  α-Tubulin

Source: [Image]
Fig. 3

A. Immunofluorescence imaging of Ishikawa PR-B cells treated with anti-PR and DAPI. Treatment groups include: -H, R5020, and RU486.

B. Western blot analysis of Ishikawa PR-B cells treated with R5020 and RU486. The bands detected are SRC-1 at 160 kDa, PR-B at 116 kDa, and α-Tubulin at 50 kDa.

C. Immunofluorescence imaging of Ishikawa PR-B cells treated with anti-SRC-1 and DAPI. Treatment groups include: -H, R5020, RU486, and MG132.

D. Western blot analysis of Ishikawa PR-B cells treated with R5020 and RU486. The bands detected are SRC-1 at 160 kDa, PR-B at 116 kDa, and α-Tubulin at 50 kDa.

E. Western blot analysis of Ishikawa PR-B parental cells treated with R5020 and RU486. The bands detected are SRC-1 at 160 kDa, PR-B at 116 kDa, and α-Tubulin at 50 kDa.

F. Western blot analysis of Ishikawa PR-B cells treated with R5020, RU486, and MG132 at different time points. The bands detected are SRC-1 at 160 kDa, PR-B at 116 kDa, and α-Tubulin at 50 kDa.
Fig. 4

A

SRC-1

\[ \Delta \text{(PEST)} \]
\[ \Delta \text{(2-16)} \]
\[ \Delta \text{(bHLH)} \]
\[ \Delta \text{(41-136)} \]

B

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<th>( \Delta \text{(bHLH)} )</th>
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<td>50 kDa</td>
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C

\[ \Delta \text{(PEST)} \] anti-S7/Rpt1 Merge DAPI / Merge

\[ \Delta \text{(bHLH)} \] anti-S7/Rpt1 Merge DAPI / Merge

D

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Fig. 5

A

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<td>α-Tubulin</td>
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Bar graph showing SRC-1/α-tubulin (AU) for R5020 treatment with SRC-1, Δ(PEST), and Δ(bHLH) conditions.

B

IP: anti SRC-1

WB: anti SRC-1

WB: anti PR

1% input

IP: anti PR

WB: anti SRC-1

WB: anti PR
Fig. 7

A) Luciferase activity (AU)

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B) Luciferase activity (AU)

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C) Relative amphiregulin expression (mRNA/18SrRNA)

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