The Heterodimeric TWIST1-E12 Complex Drives the Oncogenic Potential of TWIST1 in Human Mammary Epithelial Cells

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Abstract

The TWIST1 embryonic transcription factor displays biphasic functions during the course of carcinogenesis. It facilitates the escape of cells from oncogene-induced fail-safe programs (senescence, apoptosis) and their consequent neoplastic transformation. Additionally, it promotes the epithelial-to-mesenchymal transition and the initiation of the metastatic spread of cancer cells. Interestingly, cancer cells recurrently remain dependent on TWIST1 for their survival and/or proliferation, making TWIST1 their Achilles’ heel. TWIST1 has been reported to form either homodimeric or heterodimeric complexes mainly in association with the E bHLH class I proteins. These complexes display distinct, sometimes even antagonistic, functions during development and unequal prometastatic functions in prostate cancer cells. Using a tethered dimer strategy, we successively assessed the ability of TWIST1 dimers to cooperate with an activated version of RAS in human mammary epithelial cell transformation, to provide mice with the ability to spontaneously develop breast tumors, and lastly to maintain a senescence program at a latent state in several breast cancer cell lines. We demonstrate that the TWIST1-E12 complex, unlike the homodimer, is an oncogenic form of TWIST1 in mammary epithelial cells and that efficient binding of both partners is a prerequisite for its activity. The detection of the heterodimer in human premalignant lesions by a proximity ligation assay, at a stage preceding the initiation of the metastatic cascade, is coherent with such an oncogenic function. TWIST1-E protein heterodimeric complexes may thus constitute the main active forms of TWIST1 with regard to senescence inhibition over the time course of breast tumorigenesis.

Introduction

The TWIST1 gene is aberrantly reactivated in a large set of solid cancer types, including a variety of carcinomas, melanomas, sarcomas, and neuroblastomas [1]. As a reminiscence of its embryonic functions, this transcription factor was originally shown to promote the metastatic dissemination of cancer cells through its ability to trigger an epithelial-to-mesenchymal transition [2]. We and others additionally highlighted its ability to alleviate the induction of fail-safe programs (senescence, apoptosis) in response to oncogenic activa-...
tions, thereby cooperating with mitogenic oncoproteins in promoting cell neoplastic transformation in vitro and tumor initiation in vivo [3–9]. Furthermore, cancer cells often remained dependent on TWIST1 functions to protect them from latent senescence or apoptosis [3,4,8–11]. Its oncogenic properties are mostly related to its ability to downmodulate the activity of p53 [6,9,11,12] and to dampen the expression of numerous cyclin-kinase inhibitors (p15INK4B, p16INK4A, p21CIP1), the relative contribution of these activities varying according to the cellular context [3,4].

Basic-helix-loop-helix (bHLH) transcription factors were originally classified into different categories based on their expression pattern, partners, and structural features [13]. The TWIST1 protein belongs to class II, which encompasses tissue-specific bHLH, such as MyoD. These proteins heterodimerize with the ubiquitously expressed class I bHLH proteins, termed E proteins [13]. They include the two splice variants of the TCF3/E2A protein E12 and E47, the TCF4/ITF2, and the TCF12/HEB transcription factors. All TWIST1-E heterodimers have been shown to display similar functional properties [14]. The TWIST1 protein also interacts with the class II bHLH transcription factors HAND1 and HAND2 and additionally displays the property to constitute functional homodimers, as originally demonstrated in Drosophila [15]. The TWIST1 protein preferentially heterodimerizes, providing a rationale for the similar limb formation defects triggered by the enforced expression of the TWIST1-E12 heterodimer or of the TWIST1 monomer [16–18]. Nonetheless, homodimerization is privileged at low protein concentration (e.g., TWIST1 haploinsufficiency in Saethre-Chotzen patients), in the presence of ID HLH proteins which titrate E proteins, or following the phosphorylation of residues located in the helix I of TWIST1 (i.e., Thr121 and Ser123) by protein kinase A [14,19,20]. Point mutations preventing the posttranslational modification of these residues (TS121-123AA) or mimicking their constitutive phosphorylation (TS121-123ED) were demonstrated to functionally mimic the TWIST1 homodimer and heterodimer, respectively [16]. The phoshoregulation of TWIST1 was additionally shown to influence its affinity for E-boxes in a cis-element-dependent manner [20]. Posttranslational modifications of TWIST1 thus have an impact on dimer choice and on the downstream activation of targeted genes [21]. The contribution of these complexes to the embryonic TWIST1 functions has been largely explored using tethered dimers, the reliability of this approach being unquestionably established through successful in vivo complementation assays [15,16,19]. The validity of this strategy was also supported by the demonstration that the tethered TWIST1-E12 dimer binds DNA and transactivates reporter genes similarly to when TWIST1 and E12 are expressed as separated polypeptides [15,20]. Exploiting this tethered dimer strategy unveiled differential and even antagonistic properties of the TWIST1 complexes during embryonic development, demonstrating the key role of the TWIST1 partner in determining and regulating TWIST1 functions. Whether the balance between these dimers also modulates TWIST1 functions during tumorigenesis remains relatively unexplored. Enforced expression of tethered TWIST1-TWIST1 or TWIST1-E12 dimers in prostate cancer cell lines led to the conclusion that the prometastatic properties of TWIST1 are allotted to the heterodimeric TWIST1 complex [22]. Their contribution to the malignant transformation nonetheless still remains to be determined. Moreover, identifying the TWIST1 complex implicated in the escape from fail-safe program and upon which cancer cells are dependent for their proliferation and survival constitutes an essential step in the development of novel therapeutic strategies aiming at eradicating cancer cells through TWIST1 inactivation. To tackle this question, we employed this tethered dimer strategy to assess the oncogenic potentials of both TWIST dimers in vitro and in vivo.

**Material and Methods**

**DNA Constructs**

The EcoR pBabe Zeo, ER-H-RASG12V pLNCX2-Neo, and H-RASG12V phage-Puro retroviral construct and the VIMI-luciferase reporter have been previously described [23–26]. The tethered dimers were generated by polymerase chain reaction (PCR) using FLAG-human TWIST1 [3] and E12 expression constructs [27] as templates. The strategy employed to generate the tethered dimers is described in detail in the Supplementary Information section. The shRNA TWIST1 A pLKO.1 lentiviral construct was previously described [3]. The shRNA TWIST1 B pSIREN (5′-CTCTGGAGCTGGATAACTCAA-3′) retroviral construct was kindly provided by Patrice Lassus (IGMM, Montpellier). The shRNA TCF3 pLKO.1 was supplied by Sigma.

**Mouse Strains**

Animal maintenance and experiments were performed in a specific pathogen-free animal facility, “Anicam,” in accordance with the animal care guidelines of the European Union and French laws and were validated by the local Animal Ethic Committee (CECAAP). The CAG-LSL-Twist1-Twist1 or CAG-LSL-Twist1-E12 (FVB/NJ genetic background) [19] was crossed with the WAP-Cre (whey acidic protein promoter [28]) mouse strain [B6.Cg-Tg(Wap-Cre)11738 Mam, obtained from the mouse models of Human Cancers Consortium and back-crossed to the FVB/NJ genetic background]. Cohorts of multiparous female WAP-Cre; Twist1-Twist1 and WAP-Cre; Twist1-E12 mice (with two rounds of lactation) were generated. Genotyping was performed on genomic DNA from tails using the 5′-AATGGAATGGAGAGCTTGGGCGAC-3′ and 5′-CTCTGGAGCTGGATAACTCAA-3′ primer pair for the WAP-Cre transgene and 5′-GAACACGGCCCGCCAAGAATACTCG-3′ and 5′-CCCCTCTTGAATGGCAG-3′ primer pair for the Twist1 transgene as described previously [19,28]. Mice were monitored twice a week for tumor incidence. End points were based on tumor diameter (up to 1.5 cm) unless natural death occurred. Mice were euthanized by CO2 inhalation. Tumors were fixed in 10% phosphate-buffered formalin for 24 hours and embedded in paraffin. Mammary gland, lung, liver, kidney, and spleen were collected either to check for malignant lesions or to assess potential metastatic dissemination.

**Cell Culture**

HMEC-hTERT cells and derived cells were cultured as described previously [7]. The Hs578T, MDA-MB-436, MDA-MB-231, and BT-474 breast cancer cell lines and HEK293T were provided by the ATCC and cultured according to the supplier’s instructions.

**Protein Analysis and Immunohistochemistry**

Cells extracts were performed in radioimmunoprecipitation assay (RIPA) buffer [100 mM NaCl, 1% NP40, 0.1% sodium dodecyl sulfate (SDS), 50 mM Tris pH 8] supplemented with protease (Roche) and phosphatase inhibitor cocktails (Sigma) and clarified by centrifugation. Proteins were analyzed by Western blot using the monoclonal anti-TWIST1 2C1a (Abcam), anti-p21CIP1 clone SX118 (Dako), and anti-HEB A9 (Santa Cruz Biotechnology) and the polyclonal anti-TWIST2 (proteintech), anti-E2A V18, anti-ITF2 K12, anti-H-RAS C20, anti-p15INK4B C20, anti-p16INK4A H156, p-histone H3 (Ser10)-R...
(Santa Cruz Biotechnology), and anti-GAPDH Abs16 (Millipore) antibodies and horseradish peroxidase–conjugated secondary antibodies (Dako). Antigen-antibody complexes were revealed with the Western-blotting Luminol reagent (Santa Cruz Biotechnology).

Analysis of TWIST1 by immunohistochemistry in mouse breast carcinomas was performed as described previously [7] using the monoclonal anti-TWIST1 2C1a (Abcam).

**Gene Expression Analysis**

RNA preparation and reverse transcription were performed as described previously [29]. Real-time PCR intron-spanning primers were designed with the primer3 software. The HPR71 housekeeping gene was used for normalization. The combinations of primers used are listed in the Supplementary Information section.

**Retroviral Infections, SA-β-Galactosidase, Growth Curves, and Soft-Ag Colony Assays**

Enforced expression of H-RAS(G12V) and/or TWIST1 proteins was performed through retroviral infections as described previously [29]. Briefly, cells were “murinized” by expressing the ecotropic receptor [24,30] before being infected with retroviral expression constructs 48 hours later. Selection was initiated 24 hours post–second infection with puromycin (0.5 μg/ml) or neomycin (100 μg/ml).

shRNA TWIST1 lentiviral particles were generated through the co-transfection of 293T cells with pLKO.1, pCMV ΔR8.91 (gag-pop-Tat-Rev) [31], and phCMVG-VSVG (env) [32] expression constructs, and shRNA TWIST1 retroviral particles were generated through the transfection of HEK293GP cells with pSIREN and phCMVG-VSVG (env) expression constructs using the calcium phosphate precipitation technique. TWIST1 knockdown was achieved via double retroviral (shRNA TWIST1 B pSIREN) or lentiviral (shRNA TWIST1 A pLKO.1) infections at 48-hour intervals. Over 90% of cells were infected. Consequences of TWIST1 depletion on cell survival and proliferation were assessed in the absence of selection. Activation of ER-RAS(G12V) with 4-OHT (4-OHT, 625 nM, Sigma Aldrich) was specifically performed through retroviral infections as described previously [29].

**Immunoprecipitation**

HEK293T cells were transfected with wild-type or mutant FLAG-tagged TWIST1 and/or MYC-tagged E12 expression constructs using the calcium-phosphate technique. Cells were lysed 36 hours posttransfection in a 50-mM Tris-HCl pH 7.4, 150-mM NaCl, 1-mM EDTA, and 1% Triton buffer supplemented with protease inhibitors (Roche), cleared by centrifugation, and incubated with an anti-FLAG M2 resin (Sigma-Aldrich). After intensive washing to eliminate nonspecific binding, the resin was resuspended in Laemmli buffer and boiled for 3 minutes. After elimination of the beads by centrifugation, samples were reduced by adding β-mercaptoethanol and separated on SDS–polyacrylamide gel electrophoresis. Proteins were analyzed with the murine monoclonal anti-TWIST1 C2a antibody (Abcam) and the rabbit polyclonal V18 E2A antibody (Santa Cruz Biotechnology).

Immunoprecipitation of endogenous E12 protein by ectopically expressed FLAG-TWIST1 was performed in HMEC-hTERT cells. A total of 2 × 10^6 cells were resuspended in an extraction buffer (20 mM Tris pH 8, 125 mM NaCl, 1 mM EDTA, 0.5% NP40) supplemented with protease inhibitors (Roche), sonicated, and cleared by centrifugation. Protein extracts were diluted in a final 20-mM Tris pH 8 buffer containing 125 mM NaCl, 1 mM EDTA, NP40 0.125%, and glycerol 10% and successively incubated in the presence of 4 μg of M2 α-FLAG antibody (Sigma) for 4 hours at 4°C and protein-A sepharose beads (protein A sepharose 4B, Invitrogen) for 1 hour. After intensive washing in a 20-mM Tris pH 8 buffer containing 125 mM NaCl, 1 mM EDTA, NP40 0.1%, and glycerol 10%, beads were resuspended in Laemmli buffer (in absence of β-mercaptoethanol) boiled at 95°C for 3 minutes. After elimination of the beads by centrifugation, samples were reduced by adding β-mercaptoethanol and separated on SDS–polyacrylamide gel electrophoresis.

**Proximity Ligation Assay**

Human ductal in situ carcinoma (DCIS) samples were obtained through the Biological Resource Center of the Centre Léon Bérard with the agreement of the ethical review board of the Centre Léon Bérard. Samples were used with the patient’s written informed consent. The present study was approved by the ethical review board of the Centre Léon Bérard. DCIS samples were selected on the basis of TWIST1 protein detection, as previously reported [7].

After deparaffinization and rehydration, tissue sections were boiled in a 10-mM pH 6 citrate buffer for 40 minutes. After saturating unspecific binding sites for 20 minutes with a 1% BSA PBS buffer, sections were incubated overnight with the mouse monoclonal anti-TWIST1 Twist2C1a (Abcam) and the rabbit polyclonal anti-E2A N-649 (Santa Cruz) antibodies in a 0.5% BSA in PBS buffer. Incubation with PLA probes, ligation, and amplification steps were performed according to the manufacturer instructions (Duolink In Situ kit, Sigma). The specificity of the signal was confirmed using paraffin-embedded MDA-MB-436, Hs578T (TWIST1 positive, E12 positive), and HMEC-hTERT (TWIST1 negative, E12 positive) mammary cells in which TWIST1 or TCF3 (the two E2A protein variants E12 and E47 encoding gene) were specifically knocked down through RNA interference. Number of dots per cell was assessed using the ImageJ software.

**Results**

**Design of Wild-Type and Mutant Tethered TWIST1 Dimers**

Because both HAND1 and HAND2 genes were found to be transcriptionally inactive in breast cancer cells and because the TCF3/
E2A proteins are the predominant E proteins expressed in the breast cancer cell lines employed in this study, as assessed by Western blot (Figure S1A), the experiments were performed either with homodimeric (TWIST1-TWIST1, hereafter named T-T) or heterodimeric (TWIST1-E2A E12, hereafter named T-E) tethered dimers. Following previous studies [15,19,20,22], the E2A splice variant E12 was employed as the model for TCF3/E2A proteins. Tethered dimers were generated by linking the two partners with a flexible glycine-serine polylinker, also used in previous studies [15,19,20,22], and the TWIST1 protein was tagged at the N-terminus with a single FLAG peptide, a position that we previously confirmed does not interfere with TWIST1 oncogenic functions [3] (Figure S1B). The production and correct nuclear localization of the resulting tethered dimers were validated (Figure S1C). Noticeably, C-terminal truncated subproducts (revealed with both anti-TWIST1 and anti-FLAG antibodies) were detected. Very similar subproducts were revealed with the functional murine MYC-tagged T-T and MYC-tagged T-E forced dimers generated in the laboratory of D. Spicer [14] (Figure S1D). In particular, the detection with the T-T dimer of a predominant subproduct with a molecular size close to the monomer likely reflects a difference in the accessibility of a consensus caspase 3/7 cleavage site present in the C-terminal of TWIST1 within the two tethered dimers. Cleavage of TWIST1 by caspase 3 was originally reported in apoptotic cells and was quickly followed by its proteasome-mediated degradation [34]. It is, therefore, very likely that the cleavage occurs during protein extraction.

As the study progressed, additional mutants were generated including the T R154P-E heterodimer (hereafter named T RP-E) to confirm the need of the interaction between both partners to generate an active tethered heterodimer. The R154P mutation was previously reported to disrupt the TWIST1-E2A protein interaction [35], and this was herein confirmed by performing a co-immunoprecipitation assay (Figure S2A). Insertion of this mutation into the T-E tethered dimer (T RP-E) did not have an impact on its nuclear sublocalization (Figure S2B) but, as anticipated, abrogated the ability of the fusion protein to activate the TWIST1-targeted vimentin (VIM) gene in a reporter assay (Figure S2C). A TWIST1 K145E-TWIST1 K145E (T KE-T KE) tethered dimer was also generated as an inactive version of the T-T tethered dimer. The Lys145 residue plays a determining role in stabilizing TWIST1 complexes on the DNA by contributing to the establishment of the interhelical loops and by directly interacting with oxygen atoms of DNA bases [36,37]. The K145E mutation thus affects the global structure of TWIST1 complexes and abrogates TWIST1 DNA binding properties [38].

**The TWIST1 Dimers Unequally Alleviate Oncogene-Induced Senescence in Human Mammary Epithelial Cells**

To compare the ability of both TWIST1 dimers in preventing oncogene-induced senescence (OIS), immortalized and nontransformed human mammary epithelial cells (HMEC-hTERT) were sequentially infected with retroviral vectors encoding either for an inducible activated version of HRAS (ER-H-RASG12V) or for TWIST1 (monomer T, T-T or T-E dimer) (Figure 1A). Activation of the

![Figure 1.](image)

**Figure 1.** Homo- and heterodimeric TWIST1 complexes unequally avoid RAS-induced senescence in immortalized human mammary epithelial cells. (A) HMEC-hTERT cells were sequentially infected with the inducible ER-H-RASG12V fusion protein and with monomeric TWIST1, the homodimeric TWIST1-TWIST1 tethered dimer, or the wild-type (wt) or mutant (RP) TWIST1-E12 heterodimeric tethered dimer. (B) In the absence of TWIST1, the activation of the ER-H-RASG12V protein, using 4-OHT, was associated with the induction of senescence as shown in the SA-β-galactosidase assay and the accumulation of the cyclin-kinase inhibitors p15INK4B (p15) and p21CIP1 (p21) as assessed by Western blot. (C and D) Five days postactivation of ER-H-RASG12V using 4-OHT, cell proliferation was assessed through a crystal violet staining (C), and p21CIP1 (p21), p15INK4B and TWIST1 proteins were analyzed by Western blot (D). Relative signal quantification with respect to MOCK cells is indicated.
ER-H-RAS\textsuperscript{G12V} fusion protein by 4-hydroxyl-tamoxifen (+4-OHT) triggered a senescence program, as revealed by the permanent proliferation arrest and the detection of a SA-β-galactosidase activity (Figure 1B). While the CDKN2A-ARF locus is methylated in HMEC-hTERT cells [39–41], commitment into senescence was associated with an accumulation of p15\textsuperscript{INK4B} (p15) and p21\textsuperscript{CIP1/WAF1} (p21) (Figure 1B). In line with our previous observations [3], enforced expression of the TWIST1 monomer successfully prevented the activation of cyclin-kinase inhibitors and sustained cell proliferation, as assessed in a crystal violet coloration assay (Figure 1, C and D). The heterodimer turned out to be functionally active in this assay and the homodimer inactive (Figure 1, C and D), suggesting that the dimerization between TWIST1 and E protein partners is a prerequisite to carry out this function. To support this hypothesis, we observed that the disruption of the partner interaction through the insertion of the R154P point mutation in TWIST1 abolished T–E activity (Figure 1, C and D). Collectively, this assay demonstrated that both monomeric and tethered heterodimers similarly prevent OIS, supporting the concept that ectopically expressed TWIST1 can form heterodimers with endogenous E proteins [17, 18]. Indeed, ectopic TWIST1 was confirmed to immunoprecipitate the endogenous E12 protein (Figure S3). Of note, in light of the absence of activity of the TWIST1 homodimeric complex, the functionality of the fusion protein was further controlled by assessing its ability to transactivate a set of TWIST1 targeted genes, selected on the basis of previous studies [7, 42]. As shown in Figure S4, the homodimer was confirmed to be functional. As an internal control, the insertion of the K145E mutation (T KE–T KE tethered dimer), described to abolish the TWIST1 DNA binding capability [38], annihilated its transactivation potential.

To strengthen the differences observed between the two TWIST1 complexes, we next assessed their ability to cooperate with the H-RAS\textsuperscript{G12V} protein in promoting the neoplastic transformation of HMEC-hTERT cells. To this end, cells were sequentially infected with TWIST1 and constitutive H-RAS\textsuperscript{G12V} retroviral expression constructs. Whereas the sole expression of the TWIST1 tethered dimers failed to transform cells, concomitant production of H-RAS\textsuperscript{G12V} and the active heterodimeric complex successfully transformed cells, as assessed in a colony formation assay (Figure 2, A and B). Neither the homodimeric T–T complex nor the inactive form of the T–E complex (T RP–E) was found to be functional in this second assay.

The enforced Expression of the T–E Tethered Dimer Induces the Spontaneous Development of Breast Carcinoma in Mice

Next, we validated the different oncogenic activities observed between the two TWIST1 complexes in vivo. We previously reported that the combined production of either the murine monomeric TWIST1 or the...
T-E fusion protein with an activated version of K-RAS (K-RASG12D) in luminal committed epithelial cells induced breast tumor development in the first 5 months with a complete penetrance [7]. However, not a single one of the three mice examined, coexpressing the homodimer T-T and K-RASG12D, developed breast tumors within the same period of time, suggesting that the homodimer failed to cooperate with RAS in promoting tumor initiation (G.W. Hinkal, unpublished data). To strengthen this result, we examined whether the ectopic expression of T-T or T-E in the absence of an oncogenic insult was sufficient to promote breast carcinogenesis, with an anticipated lower penetrance and extended latency to accumulate requested secondary events. T-T or T-E production was enforced in luminal committed cells through the generation of WAP: Twist1-TWIST1 and WAP: Twist1-E12 mice (producing the murine T-T or T-E fusion proteins under the control of the whey acidic protein promoter) by crossing either the CAG-LSL-Twist1-TWIST1 or CAG-LSL-Twist1-E12 mouse strain with the WAP-Cre mouse strain [19,28]. Full activation of the WAP promoter was assured by two consecutive lactations. As shown in Figure 2, C-E, T-E-producing mice were found to develop breast carcinomas with a higher frequency (37.9%, n = 29) and a shorter latency than T-T-producing mice (18.1%, n = 14) and control littermates (6.2%, n = 16). Tumor phenotypes were found to be heterogeneous and classified [43] as (i) adenosquamous, (ii) metastatic spindle cell, (iii) papillary and cribriform, (iv) solid and cribriform, and (v) adenosquamous carcinomas (Table S1). No secondary tumor site was detected. Collectively, these observations support our conclusion that the heterodimeric complexes are the oncogenic forms of TWIST1 in mammmary epithelial cells.

The Heterodimeric TWIST1-E12 Complex Is Detected in Human Breast Premalignant Lesions

If the assumption that the heterodimeric TWIST1 complex plays a role in malignant transformation, breast cancer cell survival, and proliferation is correct, we would expect that this complex would already be detectable in DCIS before the initiation of the metastatic cascade. We thus assessed TE complex formation through a proximity ligation assay (PLA), selecting cases previously defined as TWIST1-positive or -negative by immunohistochemistry [7]. Experimental conditions were established in mammary carcinoma cell lines, and the specificity of the signal was confirmed by turning down the expression of either the endogenous TWIST1 or TCF3 (the E12/E47 proteins-encoding gene) through RNA interference. Depletion in the TWIST1 and E2A proteins significantly reduced the number of dots detectable in the TWIST1-positive MDA-MB436 (8.8 ± 1.1 dots/cell to 4.9 ± 1.0 and 1.6 ± 1.5 dots/cell, respectively) and Hs578T (9.8 ± 3.1 dots/cell to 3.1 ± 1.9 and 2.2 ± 1.7 dots/cell, respectively), whereas the number of dots/cell remained <1 in TWIST1-negative HMEC-hTERT cells (Figure S5). As shown in Figure 3, the TE complex was invariably detected in all TWIST1-positive DCIS, whereas no signal was detected in the control samples, namely, TWIST1-negative DCIS and normal mammary glands. The detection of the TE complex in premalignant lesions is thus compatible with its role in the inhibition of a fail-safe program.

The TWIST1-E12 Heterodimeric Complex, Unlike the Homodimeric TWIST1 Complex, Complements the Loss of the Endogenous TWIST1 Protein in Sustaining Mammary Epithelial Cancer Cell Proliferation

By lowering the activity of oncosuppressive pathways and of the cell cycle control machinery, the TWIST1 protein not only favors the cell neoplastic transformation but also protects cancer cells from latent OIS and/or apoptosis [3–5,8–10]. To assess the contribution of TWIST1 dimers in these secondary activities, we selected a shRNA targeting the 3′-untranslated sequences of the endogenous TWIST1 RNA, which are absent from the heterodimer cDNAs, to inhibit endogenous TWIST1 expression and performed complementation assays with the tethered dimers. We confirmed that this shRNA (shRNA TWIST1 B), similarly to the shRNA used in our previous studies (shRNA TWIST1A [3]), induced a proliferation arrest (as assessed by a crystal violet coloration assay and the amount of phospho-Ser10 histone H3) and triggered a senescence program, as revealed by the detection of a SA-β-galactosidase activity (Figure 4), in all three breast cell lines tested (basal-B/claudin-low MDA-MB-436 and Hs578T cell lines and the luminal B BT-474 cell line). The permanent growth arrest was invariably associated with an accumulation of p15INK4B (p15) and p21CIP1 (p21) cyclin-dependent kinase inhibitors, p16INK4A (p16) remaining undetectable (Figure 4). No trace of cleaved caspases-3 or PARP fragments was detected, excluding a concomitant apoptosis induction (data not shown). To evaluate the contribution of both the homodimeric and heterodimeric TWIST1 complexes, MDA-MB-436 cells were next sequentionally infected with TWIST1 expressing constructs (using a monomeric TWIST1 cDNA lacking 3′-untranslated sequence as an internal positive control) and depleted in endogenous TWIST1 through RNA interference. As shown in Figure 5, similarly to the ectopically expressed TWIST1 monomer, the heterodimeric T-E complex avoided cyclin-kinase inhibitor accumulation and sustained cell proliferation. As anticipated, the inactive T-E variant (T RP ~E) failed to do so. The T-T homodimer also failed to complement the endogenous TWIST1 functions. Similar results were obtained in the two additional Hs578T and BT-474 cell lines (Figure S6). These data unambiguously demonstrated that the T-E heterodimeric complex is necessary to prevent OIS and to maintain the senescence program in a latent state in mammary epithelial cells.

Discussion

The embryonic transcription factor TWIST1 displays pleiotropic functions during carcinogenesis, promoting the neoplastic transformation of cells and tumor initiation, on the one hand, and the metastatic spread of cancer cells through EMT induction, on the other hand [1]. Determining whether these functions rely on the activities on various TWIST1 complexes or whether the oncogenic and prometastatic properties are conferred upon a single complex is essential to further understand the TWIST1 functions as well as to design strategies to inactivate the transcription factor. By using a tethered dimer approach, previously proven to delineate the TWIST1 complex functions in vitro as well as in vivo, we demonstrated that the TWIST1-E12 heterodimer efficiently avoided OIS in mammary epithelial cells. Furthermore, its enforced expression complements the activity of the endogenous TWIST1 protein in keeping cyclin-kinase inhibitors under a threshold compatible with breast cancer cell proliferation. In comparison, the homodimeric complex failed to do so, highlighting the pivotal role of the TWIST1 partner in regulating its oncogenic functions. To support this conclusion, the monomeric TWIST1 protein and the heterodimeric TWIST1 complex were found to efficiently cooperate with RAS in promoting breast carcinogenesis in vivo [7]. Despite the limited number of mice examined, the absence of pathology observed when the homodimeric TWIST1 complex was combined with RAS demonstrated that T-T was less efficient than T-E in promoting tumor initiation in this context (G.W. Hinkal, unpublished results). The tendency of mice,
ectopically expressing T-E in mammary epithelial cells, to spontaneously develop breast carcinoma with a higher frequency than their T-T counterparts and control littermates further supports this conclusion.

Several bHLH transcription factors are known to heterodimerize with TWIST1, including the TCF3/E2A splice variants E12 and E47, the TCF4/ITF2, the TCF12/HEB, and the HAND proteins. As the HAND genes were found to be transcriptionally inactive in the examined breast cancer cells (Figure S1), we focused on E proteins, using E12 as a prototype. Considering the degree of homology in their bHLH domains (81.8% identity with TCF3/E2A E47 splice variant, 89.1% identity with TCF4/E2-2, and 92.7% identity with TCF12/HEB), it is likely that all of these complexes display similar oncogenic properties. E2A proteins are the predominant E proteins expressed in the breast cancer cell lines employed in the present study, as assessed by Western blot (Figure S1A). Unfortunately, we cannot exclude that additional TWIST1 bHLH protein partners also contribute to its oncogenic activity. For example, the TWIST1-related TWIST2 protein is expressed in the two basal-B/claudin-low breast cancer cell lines examined (Figure S1A). Although coexpression of both transcription factors at the cellular level remains to be demonstrated, the two proteins

**Figure 3.** The heterodimeric complex is detectable in breast premalignant lesions, as assessed in a PLA. Assessment of the TWIST1-E12 heterodimeric complex in a PLA in TWIST1-positive and TWIST1-negative DCIS [7]. Healthy mammary gland was used as a control. Each single dot corresponds to the detection of a TE complex.
are able to heterodimerize [44]. It is worth noting that the heterodimeric TWIST1-E12 complex was additionally shown to drive the prometastatic activity of TWIST1 in prostate cancer cells [22], suggesting that the TWIST1-E complexes are the active forms of TWIST1 with regard to tumorigenesis, promoting both tumor initiation and progression. Using a PLA, we confirmed that the heterodimer is detectable in DCIS before the metastatic cascade is initiated (Figures 3 and S5).

Because the ectopic expression of the tethered dimer T-T did not affect breast cancer cell proliferation, it is possible that the homodimeric complex does not display transdominant negative properties, at least with regard to the functions and cellular models considered in this study. The balance between the two complexes rather spatially and temporally controls the amount of heterodimeric complex and, thereby, the oncogenic and prometastatic properties of TWIST1. To support

Figure 4. Depletion in TWIST1 reactivates latent OIS in breast cancer cells. Breast cancer cells (A, MDA-MB-436; B, Hs578T; C, BT-474) were depleted in TWIST1 through RNA interference (shRNA TWIST1 A or B). (Left panels) Two days postinfection, cells were seeded at low density and stained after 3 days with crystal violet or assessed for their SA-β-galactosidase activity. (Right upper panels) Analysis of the level of the endogenous TWIST1 transcripts by qRT-PCR. Levels were expressed relative to the housekeeping HPRT1 gene transcript and were normalized against uninfected cells. Mean, SD and Student's t test of one experiment performed in triplicate are shown. (Right lower panels) Analysis of Ser10-histone H3, p15INK4B (p15), p16INK4A (p16), p21CIP1 (p21), and TWIST1 by Western blotting (relative signal quantification with respect to shRNA control infected cells is indicated). “−” means below the threshold.
this hypothesis, ID HLH proteins which tilt the balance in favor of the TWIST1 homodimer were shown to suppress the TWIST1-driven invasive properties and to favor secondary site colonization through the induction of a mesenchymal-to-epithelial transition [45,46]. Interestingly, in both Caenorhabditis elegans and mouse development, respectively, the HLH8 (the C. elegans TWIST ortholog) and HAND1 homodimeric complexes were found to play a key role, and the heterodimers’ functions were restricted to that of counterbalancing their activity [47,48]. The relative contribution of bHLH complexes during embryonic development and senescence inhibition over the time course of breast carcinogenesis may thus be inverted. Nonetheless, the ability of the homodimer to modulate downstream targeted genes suggests that it might also display unrelated and yet-uncharacterized protumoral properties.

In numerous tumor models, cancer cells were shown to remain dependent on TWIST1 to sustain proliferation [3,4,8–11]. We now extend this observation to EMT-committed basal-B breast cancer cells (such as MDA-MB-436 and Hs578T), which express a large spectrum of EMT inducers including ZEB1 and TWIST2. ZEB1, similarly to TWIST1, has been demonstrated to downregulate the expression of INK4B and CIP1 and to prevent EGFR-driven OIS in lung cancer cells [49,50]. TWIST1 and TWIST2 were also shown to similarly downregulate RB and p53 pathways [3]. The observed lack of complementation between the proteins in these breast cancer cells either suggests cell type-specific regulations (e.g., posttranslational regulations) or unveils yet-identified TWIST1-dependent properties. The key regulators leading to the specificity of the genetic program driven by TWIST1 complexes remain to be defined. Of note, a tandem E-box separated by a 5-nucleotide spacer was recently shown to be specifically recognized by the TWIST1-E2A complexes (similarly with both E2A splice variants), a tetrameric complex formed and stabilized through the interaction between the α-helices of the WR domain of TWIST1 proteins [51]. Enforced expression of the T-E heterodimer in HMEC-hTERT cells affected the expression of a limited number of genes (data not shown), suggesting that the tethered dimer strategy avoids titration of bHLH factors. This experimental approach may lead the way to defining the TWIST1-E complex downstream targeted genes and to exploring their contribution to the oncogenic activity of TWIST1.
Funding
This work was supported by the Labellisation program of the Ligue Nationale contre le Cancer and by institutional grants from the LabEX DEVeCAn (ANR-10-LABX-61) and from the LyRiC (Lyon Recherche Intégrée en Cancérologie, Institut National contre le Cancer, INCa-4664). L. J., G. R., and C. B. were recipients of fellowships from the Ligue Nationale contre le Cancer and the Foundation ARC pour la Recherche sur le Cancer.

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Acknowledgements
We are grateful to the Biological Resource Center of the Centre Léon Bérard for providing the cohort of human breast samples. We are thankful to Prof. Cornelis Murre (University of California, San Diego, USA) for providing the human E12 expression construct, Prof. William C. Hahn for providing the EcoR retroviral expression construct (Harvard University, USA), Dr. Patrice Lassus (IGMM, Montpellier, France) for providing the human E12 expression construct, Prof. William Weis (Labellisation program of the Ligue Nationale contre le Cancer, INCa-4664). L. J., G. R., and C. B. were recipients of fellowships from the Ligue Nationale contre le Cancer and the Foundation ARC pour la Recherche sur le Cancer.

Appendix A. Supplementary data
Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.neop.2016.03.007.

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