TWIST1 SILENCES FOXA1 EXPRESSION TO PROMOTE BREAST CANCER PROGRESSION

A Dissertation

by

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Submitted to the Office of Graduate and Professional Studies of Texas A&M University in partial fulfillment of the requirements for the degree of

DOCTOR OF PHILOSOPHY

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December 2016

Major Subject: Medical Sciences

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ABSTRACT

Twist1 is a basic helix-loop-helix transcription factor family that serves as one of the master regulators promoting epithelial-mesenchymal transition (EMT). Twist1 has been shown in many studies to promote EMT, stemness, invasiveness and metastasis in multiple cancers, including breast cancer. However, the genetic role of Twist1 in spontaneous breast cancer has not been investigated, and it is also unknown whether Twist1 is required for cell invasion and metastasis of spontaneously developed cancer. In this study, using a new TVA/RCIP mouse model, we disrupted Twist1 in certain mammary luminal epithelial cells and specifically induced tumors from these cells. We found that tumor cell-specific knockout of Twist1 diminished lung metastasis without affecting primary tumor initiation or growth. The diminished lung metastasis is accompanied with the decreased EMT, angiogenesis and circulating tumor cells caused by Twist1 knockout. Twist1 expression was positively associated with late-stage tumors and negatively associated with ERα and Foxa1 expression in mouse tumors. We then identified Foxa1 as a novel direct target of Twist1 in human breast cancer. We further found that Twist1 inhibits Foxa1 expression through direct binding to its proximal promoter region and recruiting Mi2/nucleosome remodeling and deacetylase (Mi2/NuRD) transcriptional repressor complex in human breast cancer cells. Moreover, Twist1 also diminished transcriptional activator AP1 binding to Foxa1 promoter. Twist1 mediated Foxa1 down-regulation is essential for promoting breast cancer migration, invasion and metastasis. Restored Foxa1 expression significantly inhibits Twist1
dependent cell migration and invasion capability of MCF7 cells through inhibiting integrin $\alpha_5, \beta_1$ and MMP9 expression. Importantly, $\text{Twist}^\text{high} \text{Foxa}^\text{low}$ correlates with the poorest prognosis in breast cancer patients. These findings highlight Twist1 as a key player in promoting breast cancer progression to a more malignant phenotype and a potential therapeutic target.
DEDICATION

I would like to dedicate this dissertation to my father Minqiang Xu, my mother Xiaomei Lan, my wife Jingwei Zhu and my son Jeremy Xu.
ACKNOWLEDGEMENTS

I would like to thank my mentor, Dr. Jianming Xu, and my committee members, Dr. Fen Wang, Dr. Jianming Xu, Dr. Wallace L. McKeehan, Dr. Peter J.A. Davies, and Dr. Yi Li, for their guidance and support throughout the course of this research. Dr. Xu has given me valuable suggestions guiding my thesis study in the lab. He is open-minded and gives me a lot of freedom in the research. More importantly, he trained me to become an independent and inspired researcher in biomedical science. I also want to thank the other committee members for sharing their precious time and inspiring suggestions in the meetings.

My gratitude also goes to the colleagues in Xu lab. I would like to thank the former visiting student Zhen Feng for initiating the RCAS-TVA system in the lab. I would like to thank Dr. Yan Xu for collaboration and contribution to the mouse model study in the project. I would like to thank Dr. Li Qin for her tremendous help in the discussion of data and techniques. I would like to thank Dr. Tong Sun, Dr. Hongmei Wu, Dr. Zhihui Yang for their valuable input in the project. I would like to thank Dr. Dongkee Lee for his help with improving immunostaining technique. I would like to thank Lan Liao and Suoling Zhou for their help in providing SCID mice and maintaining the stock of reagents and materials in the lab. Finally, I want to thank other members in the lab including Dr. Xiaobin Yu, Dr. Jean Ching Yi Tien, Yonghong Liu and Jarrod Martinez for their suggestions to the study.
Thanks also go to the department faculty and staff for making my time at Texas A&M University Institute of Bioscience and Technology a great experience, especially the program coordinator Cindy Lewis, who is always patient to students even when we make stupid mistakes or ask dumb questions. Of course, I would also thank my friends in IBT, Xi Lin, Wenjiao Li, Yuan Dai, Yanqing Huang, Ji Jing et al. It is their company that makes the life full of fun and happiness.

Finally, thanks to my family for their encouragement and especially to my wife for her patience and love.
CONTRIBUTORS AND FUNDING SOURCES

Contributors

This work was supervised by a thesis committee consisting of Professor Fen Wang [Chair] and Professor Jianming Xu [co-Chair], Professors Wallace L. McKeehan, Peter J.A. Davies of Institute of Bioscience and Technology and Professor Yi Li of Baylor College of Medicine.

The data in Figure 3.1 were provided by Dr. Zhen Feng in Dr. Jianming Xu’s lab in Baylor College of Medicine. The fluorescent staining data in Figure 3.7 were done in collaboration with Dr. Dongkee Lee in Dr. Jianming Xu’s lab in Baylor College of Medicine. All other work conducted for the thesis was completed by the student independently.

Funding Sources

This work was made possible in part by NIH grants CA112403 and CA193455 and Cancer Prevention and Research Institute of Texas grants RP120732-P5 and RP150197.
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1. INTRODUCTION

1.1 Breast cancer

1.1.1 Basic characteristics of cancer

The origin of the word “CANCER” is credited to the Greek physician Hippocrates, who is considered the “Father of Western Medicine.” Hippocrates used the terms carcos and carcinoma to describe non-ulcer forming and ulcer-forming tumors. Cancer begins when cells in a part of the body start to grow out of control. There are many kinds of cancer, but they all start because of out-of-control growth of abnormal cells which acquire genetic or epigenetic alterations. In the 2 review papers by D. Hanahan and R. Weinberg\textsuperscript{1,2}, they proposed that these alterations occur in most if not all cancer cells which contribute to the 10 hall marks of cancer including: deregulating cellular energetics, evading apoptosis, self-sufficiency in growth signals, insensitivity to anti-growth signals, avoiding immune destruction, limitless replicative potential, tumor-promoting inflammation, tissue invasion and metastasis, sustained angiogenesis, genome instability and mutation. For most of the solid tumors, the tumor related mortality is due to the metastasis to distant organs, which results from the acquired capability of tumor cells to invade and metastasize. Tumor metastasis is a multi-step process defined as the ability of tumor cells to migrate from its original site to colonize in distant organs\textsuperscript{3}. Each
of these steps can be rate limiting, as blocking any of these steps could stop the entire process (Fig 1.1).

Figure 1.1. Overview of cancer metastasis steps. (From Isaiah J. Fidler, Nature Review: Cancer, 2003\textsuperscript{3})

1.1.2 Breast cancer and classification

Breast cancer is a heterogeneous disease with different clinical and pathological features.
Classification of breast cancer may help to predict prognosis and direct personalized treatment. Comprehensive molecular profiling of breast cancer using microarray-based technology\textsuperscript{4, 6} demonstrated that the morphological heterogeneity of breast cancer was reflected at the transcriptional level\textsuperscript{7, 8}. Their data show systematic variations of the expression profiles of breast cancer. These profiles could classify breast cancers into five subtypes: luminal A, luminal B, normal breast-like, ERBB2+ and basal-like breast carcinomas\textsuperscript{4, 6, 8, 9}. The basal-like subtype was characterized by high expression of keratins 5 and 17, laminin, and FABP7, whereas the ERBB2+ subtype was characterized by high expression of several genes in the ERBB2 amplicon at 17q22.24 including ERBB2 and GRB7. Normal breast-like group showed the highest expression of many genes known expressed in adipose tissue and other non-epithelial cell types. On the other hand, luminal type of tumors can be further grouped into two subgroups: luminal subtype A, with the highest expression of the ER\textalpha gene, GATA3, XBP1, TFF3, Foxa1, and LIV-1; and luminal subtype B with low to moderate expression of the luminal-specific genes including the ER\textalpha cluster\textsuperscript{4}.

More importantly, the molecular profile of tumor will not stay fixed as the progression of cancer. Several studies have shown that mouse mammary gland tumors progress from ER\textalpha positive/luminal A subtype to luminal B or ER\textalpha negative subtypes\textsuperscript{10, 11}. Similarly, loss of ER\textalpha expression is also broadly observed in human breast cancers, concomitant with hormone resistance, invasiveness and poor prognosis\textsuperscript{12}. 
Different gene expression profiles reflect variation in the cellular biology of the tumor subtypes and are associated with patients’ clinical outcome\(^8, 9\). Luminal A tumors have the most favorable prognosis, normal like tumors have an intermediate prognosis\(^4, 5, 9, 13\). On the other hand, luminal B, ERBB2-positive and basal-like tumors are associated with poor relapse-free and overall survival. Basal-like breast cancer (BLBC) is a major clinical challenge due to its aggressive nature and the lack of expression of the available drug targets including steroid hormone receptors (ER and PR) and ERBB2. BLBCs do not response to currently available targeted therapies such as tamoxifen (ER inhibitor) or trastuzumab (HER2 inhibitor) therapy. The absence of effective targeted therapies and poor response to standard chemotherapy may be associated with poor prognosis of BLBC patients. BLBC also has stem-cell-like properties and activated EMT (epithelial-mesenchymal transition) program, suggesting that EMT regulators may play critical roles in BLBC progression.

1.2 EMT

1.2.1 Basics of EMT

Epithelial and mesenchymal cells differ in various functional and phenotypic characteristics. Epithelial cells form a single layer or multilayers that are closely adjoined through specialized intercellular junctions such as tight junctions, adheren junctions, desmosomes and gap junctions\(^14\). Epithelial cells also have apical–basolateral
polarization, as the epithelial cells are all polarized in the same manner. Every epithelial cell membrane can be divided into an apical domain and a basolateral domain, with the apical surface facing either the exterior environment or the luminal space and the basolateral domain can be further divided to basal subdomain mediating cell-extracellular matrix adhesion and lateral subdomain involved in cell-cell contact. Cells or cell aggregates lacking the above criteria are defined as mesenchymal cells. Mesenchymal cells contact neighboring mesenchymal cells only focally and do not associate with a basal lamina. In 2D cell culture, mesenchymal cells are characterized by spindle shaped, fibroblast-like morphology, whereas epithelial cells grow as clusters that maintain cell–cell adhesion. Cultured mesenchymal cells tend to be highly motile. Epithelial cells are also motile and can move away from their nearest neighbors, while remaining within the epithelial layer. However, cells do not detach and move away from the epithelial layer under normal conditions\textsuperscript{15}.

While epithelial and mesenchymal cell types have long been recognized in early embryos, the conversion of epithelial cells into mesenchymal cell, which is known as epithelial-mesenchymal transition (EMT), was not defined until 1980s. Greenburg and Hay showed that when epithelial cells were cultured in 3D collagen gels, these cells elongated, detached from the explants, and migrated as individual cells, which are all belong to mesenchymal phenotypes\textsuperscript{16-18}. The transition from epithelial- to mesenchymal-cell characteristics involves a spectrum of inter- and intracellular changes. Subsequent molecular studies loosely defined EMT by three major changes in cellular phenotype\textsuperscript{19}. 
The first is morphological changes from monolayer of epithelial cells with an apical-basal polarity to dispersed, spindle-shaped phenotype. The second is differentiation marker changes from cell-cell junction proteins and cytokeratin intermediate filaments to vimentin filaments and fibronectin. The third is the functional changes associated with the conversion of relative stationary cells to motile cells that can invade through ECM. As not all three changes are always observed during EMT, acquisition of the ability to migrate and invade ECM as a single cell is then considered a functional hallmark of the EMT program.

Several \textit{in vitro} and \textit{in vivo} studies aimed to define the molecular regulation of EMT showed that EMT can be triggered by a crosstalk of extracellular signals including extracellular matrix (ECM) as well as soluble growth factors such as members of the TGFβ and fibroblast growth factor (FGF) families, epidermal growth factor (EGF) and SF/HGF (Fig 1.2). These growth factors serve as ligands for their cell surface receptors, which are activated in response to these ligands and trigger the activation of intracellular effector molecules, such as members of the small GTPase family — Ras, Rho and Rac and members of the Src tyrosine-kinase family. These effectors are responsible for the disassembly of junctional complexes and the changes in cytoskeletal organization during EMT. On the other hand, several transcriptional regulators such as zinc-finger transcription factors snail\textsuperscript{21}, slug\textsuperscript{22}, ZEB1\textsuperscript{23} and ZEB2(SIP1)\textsuperscript{24} and bHLH transcription factors E47\textsuperscript{25} and Twist1\textsuperscript{26} were reported to regulate the changes in gene-expression patterns that underlie EMT.
Figure 1.2. Overview of the molecular networks that regulate EMT. Signaling pathways that are activated by EMT regulators and a limited representation of their crosstalk are shown. RTKs (receptor tyrosine kinases); TGFβ (transforming growth factor-β); MMPs (Matrix metalloproteinases); ETαR (endothelin-A receptor); FAK (focal adhesion kinase); GSK3β (glycogen-synthase kinase-3β); H/E(Spl) (hairy/enhancer of split); ILK (integrin-linked kinase); MAPK (mitogen-activated protein kinase); NF-κB (nuclear factor-κB); PAR6 (partitioning-defective protein-6); PI3K (phosphatidylinositol3-kinase); PKB (protein kinase-B); ROS (reactive oxygen species); TAK1 (TGFβ-activated kinase-1); TGFβR (TGFβ receptor); WntR (Wnt receptor). (From Jean Paul Thiery and Jonathan P. Sleeman, Nature Reviews: Molecular Cell Biology, 2006)
1.2.2 EMT in development

During embryonic development, epithelial sheets can remodel through distinct processes and can also convert reversibly or irreversibly into mesenchymal cells through EMT. The process has been observed during a variety of tissue remodeling events, including mesoderm formation, neural crest development, heart valve development and secondary palate formation. Mesoderm formation and neural crest development occur during early embryonic development, epithelial cells undergo EMT, resulting in mesenchymal and neural crest cells that maintain the potential to further differentiate into various cell types. On the other hand, during heart valve development and secondary palate formation, EMT occurs in relatively well-differentiated epithelial cells that are destined to become defined mesenchymal cell types.

The earliest example of an EMT program participating in embryogenesis that been reported is the formation of mesoderm from the primitive ectoderm, mainly from model organisms including Drosophila and amphibian and avian embryos\textsuperscript{27}. Later studies indicate that the same basic principles also happen in mammalian embryos\textsuperscript{28} as epithelial cells lose their tight cell-cell adhesions and remain attached to neighboring cells only by sparsely distributed focal contacts. Subsequently, these cells undergo mesenchymal differentiation and migrate along the narrow extracellular space underneath the ectoderm, indicating the EMT program during gastrulation. Another example of EMT in embryogenesis is the generation of the neural crest, a defining tissue of vertebrates,
which is composed of a population of precursor cells with the ability to migrate over long distances in the embryo. The emergence of neural crest cells begins with the presence of a distinct population of cells with rounded and pleiomorphic shapes at the boundary between the neural plate and the epidermal ectoderm, the presumptive neural crest cells, which proceed to lose N-cadherin-mediated cell-cell adhesion while becoming excluded from the neural epithelium. Detailed studies in avian and mouse embryos using immune-labeling and electron microscopic revealed that disruption of the basal lamina occurs immediately before or at the onset of neural crest cell migration in the cranial regions. These observations demonstrate that neural crest cells actively invade through the basal lamina, however, not present at the neural fold in the trunk region before the onset of neural crest migration. These findings suggest that distinct types of subcellular machineries were employed to initiate EMT, invade the ECM, and migrate.

Heart valve in vertebrate embryos, which is derived from a precursor structure known as endocardial cushion, begins to form soon after the primitive linear heart tube begins to loop. Initially, the myocardial cells secrete a large amount of ECM and create endocardial cushions, and then the mesenchymal cells fill the cushion space. These mesenchymal cells were then found to derive from the endocardial cell layer by undergoing an EMT. Upon EMT-inducing stimuli, AV endocardial cells decrease N-CAM, lose cell-cell adhesion, and invade the newly deposited endocardial cushion, thereby establishing the presumptive cardiac septa and valves. Another well-studied
EMT is the formation of the secondary oral palate when researchers can trace the entire program of palate remolding with electron microscopy and immunohistochemistry by dissecting the palatal tissue and introducing it into organ cultures. As fusion of the palatal shelves at the midline is required for development of the secondary palate, when the shelves approach one another from opposite sides of the developing oral cavity, epithelial cells covering the tip of each shelf intercalate and form the medial epithelial seam. Then these medial epithelial cells undergo EMT and are integrated into the mesenchymal compartment of the palate to finish the program of palatogenesis36.

1.2.3 EMT in diseases

Under physiological condition, such as embryogenesis, EMT program plays an important role. However, as well as many other programs, similar cell changes could be recapitulated during pathological processes, such as fibrosis and cancer. Many studies have shown a remarkable similarity between the regulatory signaling pathways of EMT in physiological and pathological conditions.

Fibrosis is characterized by increased numbers of myofibroblasts that deposit interstitial ECM, and the origin for majority of these fibroblasts is thought to be tubular epithelial cells that undergo through EMT37. In animal models of kidney disease, EMT occurs in tubular epithelial cells38. Moreover, similar observations have been made in studies of human renal fibrosis39. One of the mechanisms could be inflammatory processes in
response to injury of tissue lead to increased levels of factors such as TGFβ, EGF and FGF2, which have been demonstrated to be EMT stimulators of tubular epithelial cells\textsuperscript{40}. Indeed, one study showed that in the peritoneal cavity, renal dialysis causes injury to the mesothelial lining and results in EMT of mesothelial cells and finally fibrosis through inducing TGFβ expression\textsuperscript{41}, which then has been confirmed in animal models\textsuperscript{42}.

Beside fibrosis, numerous observations showed that EMT plays an important role in the progression of multiple cancers, especially in tumor metastasis. Majority of solid tumors are carcinomas that originate from epithelial cells. When the carcinoma cells metastasize, they must lose cell-cell adhesion and acquire motility, which shares many similarities with the extensive cell migration and tissue rearrangements that occur during the various developmental events mentioned above. Indeed, this highly conserved EMT program has been implicated in giving rise to the dissemination of single carcinoma cells from primary epithelial tumors\textsuperscript{43}. Many reports that study the function of EMT regulators and transcriptional factors in cancer revealed the central role of these factors in facilitating tumor cell migration, invasion and metastasis.

Thus, based on the functional consequences and biological context, EMT is classified into 3 distinct types, which are developmental (Type I), fibrosis and wound healing (Type II), and cancer (Type III)\textsuperscript{44}. As mentioned above, these 3 types of EMT occur at different micro-environmental context and show different characteristics. On the other hand, these 3 types of EMT share some common mechanisms. Biomarkers specific to
each subtype of EMT and also common to all subtypes have been defined, as summarized in a recent review paper of EMT\textsuperscript{45} (Table 1.1).

Tumor cells metastasize to distant organs is the major reason of mortality in breast cancer patients. However, the mechanism by which epithelial tumor cells escape from the primary tumor and colonize a distant site is not entirely understood. Numerous studies in breast cancer and other cancers have postulated EMT as an important mechanism for tumor metastasis. Except for several rare subtypes such as diffuse lobular carcinoma, in most breast cancer, similar to other epithelial carcinomas, EMT rarely occurs homogenously across the whole tumor. However, the direct evidence of EMT existence is not easy to find as once the epithelial tumor cells undergo EMT, they may be phenotypically indistinguishable from fibroblasts. In addition, since most metastatic lesions exhibit an epithelial phenotype, whether EMT indeed occurs in breast tumor progression is still debatable until a transgenic mice study provides direct evidence of EMT in the local invasion of tumor cells in breast cancer\textsuperscript{46}. Furthermore, microarray analysis of breast tumors showed that certain types of breast cancer cells are primed to undergo EMT and spread, such as the basal or triple negative breast cancer subtype which exhibits an aggressive phenotype and correlates with the poorest clinical outcomes\textsuperscript{47}. Similar results were observed in breast cancer cells lines as basal B subtype breast cancer cell lines also display a mesenchymal phenotype\textsuperscript{48}. These work revealed the existence of EMT in primary breast cancer, which is supported by more and more evidence to be a reversible and transient process. This could explain why metastatic cells
at a secondary site are likely undergo a reversion EMT or MET, permitting colonization of the distant site\textsuperscript{49}.

1.2.4 EMT in breast cancer

It was hypothesized that developmental programs are reactivated during tumorigenesis and contribute to tumor progression. The fact that numerous EMT regulators in development are also inappropriately expressed in human cancer and correlate with features of EMT is a strong evidence to support the hypothesis\textsuperscript{22, 23, 26}. E-cadherin, an important caretaker of the epithelial phenotype, is down-regulated by these EMT transcriptional regulators. In both cell culture and tumor mouse models, ectopically expression of E-cadherin in certain invasive carcinoma cells can inhibit their capability to invade and metastasize. Conversely, downregulation of E-cadherin (or mutations that occur in cancer) has several important consequences that are of direct relevance to EMT and subsequent cell motility and metastasis. Decrease of E-cadherin level in carcinoma cells results in the loss of E-cadherin-dependent intercellular epithelial junctional complexes, and abolished sequestering of $\beta$-catenin in the cytoplasm, subsequently release the carcinoma cells from the neighbor cells.
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Table 1.1. Markers of EMT (From Michael Zeisberg and Eric G. Neilson, J.Clin. Invest, 2009)
Besides promoting tumor cell invasion and metastasis, EMT is reported to be involved in multiple aspects of tumor progression including cell survival, chemo-resistance, stemness and cancer cell transition into endothelial cells\textsuperscript{50} (Fig 1.3). For example, Robert A. Weinberg’s group reported that the induction of EMT in immortalized human mammary epithelial cells (HMLEs) results in the acquisition of mesenchymal traits and in the expression of stem cell markers. Moreover, stem-like cells isolated from HMLEs, mouse or human mammary glands and mammary carcinomas undergo EMT\textsuperscript{51}. They then update their study recently showing that distinct EMT programs control normal mammary stem cells and tumor-initiating cells\textsuperscript{52}. Another study showed that Six1 expands the mouse mammary epithelial stem/progenitor cell pool, which is correlated with EMT\textsuperscript{53}. The acquisition of a cancer stem cell-like phenotype enables a subset of tumor cells to self-renew and likely to ultimately metastasize to distant organs.
1.3 Twist1

1.3.1 Twist1 in development and human disease

A mutant *Drosophila* embryos failed to gastrulate normally, produced no mesoderm and died at the end of embryogenesis with a ‘twisted’ appearance\(^5\), thus the gene that is responsible for the phenotype is named “*twist*”. The human and mouse homolog of *twist*
gene are found and named *Twist1*. The *twist* gene encodes a transcription factor containing a basic helix-loop-helix (bHLH) domain\textsuperscript{55} involved in the regulation of organogenesis\textsuperscript{56, 57}. The critical roles of Twist1 protein in mesoderm associated organogenesis, especially in the morphogenesis of the cranial neural tube\textsuperscript{58}, have been well illustrated by genetic studies. In *Drosophila*, *twist* transcription is activated by Dorsal in the presumptive mesoderm and the expressed twist protein forms a steep gradient across the presumptive mesoderm-neuroectoderm border in the early embryo\textsuperscript{59, 60}. During later stage, *twist* expression decreases to relatively low levels in the mesodermal layer cells. The similar effect of Twist1 is found in mouse mesoderm development. After birth, Twist1 can be seen in adult mesenchymal cells such as muscle stem cells, referred to as adult muscle precursors in *Drosophila*, while in mouse, *Twist1* is expressed in the adult stem cells of the mesenchyme\textsuperscript{61, 62}. Previous study in our lab revealed that Twist1 is expressed in a few tissues, including fibroblasts of the mammary glands and dermal papilla cells of the hair follicles in young and adult mice. Conditional *Twist1* knockout in 6-week-old males and female mice did not influence body weight gain, heart rate, or total lean and fat components. The knockout also did not alter blood pressure in males, although it slightly reduced blood pressure in females. These results indicate that Twist1 is not essential for maintaining an overall healthy condition in young and adult mice\textsuperscript{63}. In summary, *Twist1* is highly expressed in the mesoderm-derived mesenchyme during embryonic development and is primarily expressed in relatively quiescent adult stem cells located in mesoderm-derived mesenchymal tissues postnatally\textsuperscript{64}. 

In humans, *Twist1* gene mutations are responsible for an autosomal dominant inheritance disease named Saethre-Chotzen syndrome (SCS), which is characterized by a broad spectrum of malformations including short stature, craniosynostoses, high forehead, ptosis, small ears with prominent crus, and maxillary hypoplasia with a narrow and high palate\textsuperscript{65, 66}. Non-sense mutations and mis-sense mutations at conserved residue across species within bHLH domain were found in human SCS patients. Since majority of mutations are heterozygous, leading to loss of function of Twist1 protein, it suggests that protein level is essential for the function of Twist1 in cells, which is supported by the gradient expression of Twist1 protein at the development stage in *Drosophila*. Gene ablation experiments of mouse models demonstrated that the heterozygous *Twist1* null mice\textsuperscript{58} manifest craniofacial and limb abnormalities resembling those in SCS patients. Since the last decade, numerous studies have shown that Twist1 also plays important roles in cancer progression, especially in cancer metastasis\textsuperscript{26, 67, 68}.

### 1.3.2 Twist1 promotes EMT in cancer

Yang *et al.*\textsuperscript{26} reported that increased *Twist1* expression correlates with breast cancer cell invasion and metastasis, and suppression of *Twist1* expression inhibits the cells’ capability to metastasize from the xenograft tumor cells in mammary gland to the lung. They also observed that expression of Twist1 protein in the epithelial cell lines results in partial EMT and induction of cell motility. Subsequent studies revealed that Twist1 promotes cancer cell survival, drug resistance, cancer stem-like cell number,
invadopodia formation for extracellular matrix degradation, and cancer cell transition into endothelial cells, respectively\textsuperscript{12, 69-72}. Previous studies revealed several regulators of Twist1 and Twist1 target genes in breast cancer. Ling and Arlinghaus\textsuperscript{73} observed that knockdown of STAT3 eliminated Twist1 expression, which does not affect cells proliferation but impair cell invasiveness. These observations suggest that STAT3 enhances Twist1 expression in its promotion of breast cancer progression. Another study\textsuperscript{74} showed that activation of STAT3 increased Twist1 expression while inhibition of STAT3 significantly reduced Twist1 expression in the aggressive human breast cancer cell lines. This study also showed that STAT3 binds directly to the second proximal STAT3-binding site on the human Twist1 promoter and activates Twist1 transcription. Together with another observation that Twist1 transcriptionally induces AKT2 to promote oncogenic functions\textsuperscript{75}. Cheng et al.\textsuperscript{70} proposed that STAT3, Twist1 and AKT2 form a functional signaling axis to enhance tumor invasion, which suggests an attractive target for cancer therapy. Interestingly, a recent study demonstrates that AKT1 inhibits EMT in breast cancer through Twist1 degradation\textsuperscript{76}. Our group found that the steroid receptor coactivator-1 (SRC-1) specifically promotes breast cancer metastasis without affecting primary tumor growth\textsuperscript{77} in a PyMT mouse breast cancer model. Subsequent investigation revealed that SRC-1 serves as a coactivator for the transcription factor PEA3 to enhance Twist1 expression, suggesting a molecular mechanism by which SRC-1 promotes breast cancer invasiveness and metastasis\textsuperscript{78}. A recent study showed an inverse relationship between TRIM29 and Twist1 and more importantly, TRIM29 and Twist1 inhibit each other in breast cancer cells\textsuperscript{79}. Twist1 also plays a critical role in
transducing the mechanical signal through G3BP2, which explains how matrix stiffness drives EMT and tumor metastasis\textsuperscript{80}.

Figure 1.4. Possible regulatory network of Twist1 in cancer progression. Twist1 is activated at transcriptional and post-transcriptional levels and promotes EMT, invasion, migration, metastasis, cancer stemness and chemo-resistance through multiple signaling pathways. (From Qian Qin, Young Xu, Tao He, Chunlin Qin, Jianming Xu, Cell Research, 2011\textsuperscript{64})

As a transcription factor, Twist1 could either activate or repress its target genes. SET8 is a member of methyltransferase family that specifically targets H4K20 for
monomethylation\textsuperscript{81, 82} and is implicated in regulating either activation\textsuperscript{83} or repression\textsuperscript{82} of gene transcription. Feng Yang\textsuperscript{84} and colleagues reported that SET8 physically associates with Twist1 to act as a dual epigenetic modifier of Twist1 target genes: repressing E-cadherin and activating N-cadherin. A recent study by Jian Shi\textsuperscript{85} and colleagues demonstrated that Twist1 binds to BRD4 thereby constructing an activated Twist1/BRD4/P-TEFb/RNA-Pol II complex at the WNT5A promoter and enhancer to promote invasion, cancer stem cell (CSC)-like properties, and tumorigenicity of BLBC cells. Fu et al. reported that Twist1 interacts with several components of the Mi2/nucleosome remodeling and deacetylase (Mi2/NuRD) protein complex, including metastasis-associated protein 2 (MTA2), Rb-associated protein 46 (RbAp46), Mi2 and histone deacetylase 2 (HDAC2)\textsuperscript{68}. Twist1 recruits this gene repression protein complex to the E-cadherin promoter, resulting in repression of the E-cadherin promoter activity and E-cadherin expression. Another study showed that Twist1 recruited DNA methyltransferase 3B (DNMT3B) to the ER\(\alpha\) promoter, leading to a significantly higher degree of ER\(\alpha\) promoter methylation. Furthermore, they demonstrated that Twist1 interacted with histone deacetylase 1 (HDAC1) at the ER\(\alpha\) promoter, causing histone deacetylation and chromatin condensation, further reducing ER\(\alpha\) transcript levels\textsuperscript{12}. 
Figure 1.5. Twist1 recruits Mi-2/NuRD complex to repress E-cadherin transcription in cancer progression. Twist1 forms heterodimer with E12, and physically interact with MTA2 (metastasis-associated protein 2), RbAp46 (Rb-associated protein 46) and Mi-2 in the Mi-2/NuRD transcription repressor complex. (From Qian Qin, Young Xu, Tao He, Chunlin Qin, Jianming Xu, Cell Research, 2011)

1.4 Foxa1

1.4.1 Foxa1 in development

Hepatocyte Nuclear Factor (HNF) was initially cloned and purified from mammalian liver, and then found also expressed in other organs. Later on, the HNF superfamily was
identified as sharing a highly conserved central motif with that of the *Drosophila melanogaster* forkhead protein (fkh), known to bind DNA and regulate processes necessary for early fly development. The HNF transcriptional regulators represent a unique subclass of DNA binding proteins that play a critical role in tissue-specific differentiation and organogenesis. The subsequent genome-wide analysis in multiple systems uncovered hundreds of forkhead transcription factors, over 40 of which were known to be expressed in mammalian systems that could be classified into more than 19 subclasses. To better clarify the nomenclature, the vertebrate HNF family was renamed “FOX” for Forkhead Box, with a letter showing the subclass, such as FOXA, FOXB. The FOXA family which contains 3 members (Foxa1, FOXA2 and FOXA3) plays pivotal roles in mammalian embryonic development and organogenesis.

*Foxa1* is expressed in the liver, pancreas, bladder, prostate, colon, lung as well as mammary gland in mice. Early studies showed that Foxa1 and another family member Foxa2 display overlapping roles in development and differentiation of the pancreas, liver and neurological system. On the other hand, Foxa1 alone appears to be the master regulator in tissues that dependent on sex hormone signaling, such as the breast and prostate glands. In the developing mammary gland, Foxa1 and ERα are co-expressed within luminal epithelial cells, with strong expression observed in the terminal end buds. *Foxa1* deletion results in phenotype similar to ERα null phenotype in ductal morphogenesis but not in alveologenesis. Foxa1 may also play a role in sustaining the luminal epithelium in an undifferentiated state.
1.4.2 Foxa1 in human cancers

Foxa1 and FOXA2 were reported to play various roles in multiple human cancers, including pancreatic cancer, thyroid cancer, prostate cancer and breast cancer. The relationship between Foxa1 and nuclear receptors has gained a lot of attention ever since the identification of this co-factor. Mapping of ERα specific binding sites on chromosomes 21 and 22 showed that ERα predominantly occupies enhancer-like regions near oestrogen responsive genes. And remarkably, they found forkhead motifs enriched in 56% of ERα-bound chromatin at nearby region. Further analyses revealed that Foxa1 occupies these regions even before ERα was recruited and serves as a pioneer transcription factor that facilitates an open chromatin structure in the absence of estrogen.

Several studies have shown that Foxa1 is required for optimum expression of around 50% of ERα regulated genes and estrogen-induced proliferation. Sunil Badve et al did immunohistochemistry for Foxa1 protein in breast carcinoma tissues of 438 patients and correlated Foxa1 expression with various established disease markers and with breast cancer-specific survival. They conclude that Foxa1 expression is associated with ERα positivity, the luminal subtype A, and better breast cancer specific survival. Concomitant studies showed similar result and also indicated Foxa1 is a critical determinant of ERα activity and endocrine response in breast cancer.
Foa1 is not only playing a role as ERα co-factor in breast cancer, but also has its own contribution to breast cancer progression such as repressing basal phenotype\textsuperscript{101}, and cross-talk to ERBB2 signaling pathway\textsuperscript{102}. Transient Foxa1 silencing results in a molecular profile transition toward basal phenotype and increases the in vitro aggressiveness of luminal breast cancer cells. Indeed, loss of Foxa1 may lead to growth arrest of a subpopulation of differentiated tumor cells, while the remaining cells may have greater chance to de-differentiate towards the basal phenotype. Foxa1 silencing may enrich this population, resulting in the observed shift toward the basal subtype\textsuperscript{101}. This notion is supported by several studies based on MMTV-PyMT model: Elaine Y. Lin et al reported that MMTV-PyMT transgenic breast cancer model developed at late stage are characterized by decreased or loss of ERα and PR expression\textsuperscript{10}; and Kasi McCune et al further discovered tumor transition from luminal A subtype to luminal B subtype concomitant with decreased ERα and Foxa1 expression\textsuperscript{11}. However, the underlying mechanisms of Foxa1 silencing are still unknown.

In this study, we examined the endogenous Twist1’s function in breast cancer progression using a novel RCIP-TVA system-mediated tumor cell specific knockout of Twist1. We found that specific knock out of Twist1 in mammary gland tumor cells does not influence tumor initiation or growth but significantly decreased tumor EMT and metastasis. We also identified Foxa1 as a direct target of Twist1 in breast cancer. Twist1 inhibits Foxa1 expression at transcriptional level and promotes breast cancer cell basal-like phenotype, invasiveness and metastasis.
2. MATERIAL AND METHODS

2.1 The TVA/RCIP system for induction of mouse mammary gland tumorigenesis with tumor cell-specific Twist1 knockout

The Cre DNA fragment with a Not I and an Asc I sites was amplified by PCR, and a small IRES DNA fragment with an Asc I and a Pac I sites was chemically synthesized. These two fragments were cloned to the upstream of the PyMT-HA coding sequence through a Not I and a Pac I sites in the RCAS-PyMT avian-viral vector. The constructed RCAS-Cre-IRES-PyMT-HA (RCIP) vector was transfected into DF1 cells to produce RCIP avian virus as described previously. The MMTV-TVA transgenic mouse line was described previously. MMTV-TVA mice were crossbred with the ROSA26R Cre reporter mice to generate MMTV-TVA;ROSA26R bi-genic mice. MMTV-TVA mice were also crossbred with Twist1F/F mice obtained from Mutant Mouse Regional Resource Centers (016842-UNC) to generate MMTV-TVA control (MMTV-TVA;Twist1+/+) and MMTV-TVA;Twist1F/F mice. All mice used in experiments were backcrossed at least for two generations in FVB strain background. PCR-based genotype analyses were carried out as described previously for the WT, floxed and deleted Twist1 alleles, the ROSA26R Cre reporter allele as well as the MMTV-TVA transgene. PCR primers were listed in Supplemental Table S1. For inducing mammary tumorigenesis, RCIP virus was intraductally introduced into the 4th mammary gland(s) of mice as described previously. The injected mice were closely examined for mammary tumor initiation by palpation, tumor growth by...
estimating tumor volume, and distant metastasis by eye and microscopic examinations as described previously. Mice with mammary tumors were sacrificed 12 or 18 weeks after viral injection. Mice were euthanized according to the NIH guidelines. The animal protocols were approved by the IACUC of Baylor College of Medicine.

2.2 Western blot

Western blot was carried out as described previously. Briefly, 20 μg of protein in cell lysate was separated in SDS-PAGE gel and blotted onto a nitrocellulose membrane. After being blocked with phosphate-buffered saline containing 5% bovine serum albumin (BSA), the membrane was incubated with a primary antibody overnight at 4°C. These primary antibodies included antibodies against Twist1 (ab50887), Foxa1 (ab23738), c-FOS (ab53036, Abcam, Cambridge, MA), E-cadherin (610182, BD Bioscience, San Jose, CA), VIMENTIN (5741), c-JUN (9165), HA (3724, Cell Signaling, Danvers, MA) and β-actin (Sigma, St Louis, MO). Then, the membrane was washed 3 times in phosphate-buffered saline with Tween-20 (PBST) and incubated with horseradish peroxidase (HRP) conjugated secondary antibody (Bio-rad, Hercules, CA) for 1 hour. In certain assays, the TrueBlot Western Blot Kit (eBioscience, San Diego, CA) with the secondary antibody was used to eliminate IgG bands according to the manufacture’s instruction. Finally, the membrane was incubated with the SuperSignal West Pico Chemiluminescent Substrate (30479; Thermo Scientific, Rockford, IL) and imaged by exposing to X-ray films.
Western blot was carried out as described previously \(^{63,111}\). Briefly, 20 \(\mu\text{g}\) of protein in cell lysate was separated in SDS-PAGE gel and blotted onto a nitrocellulose membrane. After being blocked with phosphate-buffered saline containing 5% bovine serum albumin (BSA), the membrane was incubated with a primary antibody overnight at 4\(^\circ\)C. These primary antibodies included antibodies against Twist1 (ab50887), Foxa1 (ab23738), c-FOS (ab53036, Abcam, Cambridge, MA), E-cadherin (610182, BD Bioscience, San Jose, CA), VIMENTIN (5741), c-JUN (9165), HA (3724, Cell Signaling, Danvers, MA) and \(\beta\)-actin (Sigma, St Louis, MO). Then, the membrane was washed 3 times in phosphate-buffered saline with Tween-20 (PBST) and incubated with horseradish peroxidase (HRP) conjugated secondary antibody (Bio-rad, Hercules, CA) for 1 hour. In certain assays, the TrueBlot Western Blot Kit (eBioscience, San Diego, CA) with the secondary antibody was used to eliminate IgG bands according to the manufacture’s instruction. Finally, the membrane was incubated with the SuperSignal West Pico Chemiluminescent Substrate (30479; Thermo Scientific, Rockford, IL) and imaged by exposing to X-ray films.

2.3 X-gal staining

The RCIP virus-infected inquinal mammary glands of MMTV-TVA;ROSA26R mice were collected on day 3, 21 or 48 after the virus was introduced. The mammary glands were slightly fixed and stained by X-gal as described previously \(^{112}\). The X-gal-stained mammary glands were further fixed in 4% paraformaldehyde (PFA), washed in
phosphate-buffered saline (PBS), dehydrated in ethanol and embedded in paraffin blocks as described. Five-μm thick tissue sections were prepared and used for H&E staining.

2.4 Cell culture

DF1 chicken fibroblast cells, HEK293T human embryonic kidney cells and MCF7 human breast cancer cells were cultured in MDEM medium containing 10% of fetal bovine serum (FBS). BT549 cells were cultured in RPMI-1640 medium with 10% of FBS. SUM1315 cells were cultured in Hyclone Nutrient Mixture with 5% of FBS, 5 μg/ml of insulin and 10 ng/ml of epidermal growth factor. All cells were cultured at 37°C in a tissue culture incubator supplied with 5% CO₂.

2.5 Circulating Tumor Cell (CTC) culture

About one milliliter of blood sample was collected from each mouse with mammary gland tumors. CTCs were enriched by removing red blood cells using RBC lysis buffer (420301, Biolegend, San Diego, CA) according to the manufacture’s instruction. CTCs were cultured in MDEM medium containing 10% of fetal bovine serum (FBS) at 37°C in a tissue culture incubator supplied with 5% CO₂ for 2 weeks. CTCs that form colonies which contain more than 10 cells were counted and colony numbers from two groups of mice were analyzed with student T-test.
2.6 H&E staining, immunohistochemistry (IHC) and immunohistofluorescence (IHF)

Mammary tumors and/or lungs were dissected from euthanized MMTV-TVA;\textit{Twist1}^{F/F} and MMTV-TVA mice as well as SCID mice injected with MCF7\textsuperscript{Ctrl}, MCF7\textit{Twist1} and MCF7\textit{Twist1+Foxa1} cells. Isolated tissues were fixed overnight at 4\textdegree C in 4\% PFA, washed in PBS, dehydrated in ethanol and embedded in paraffin blocks. Five-\textmu m thick tissue sections were prepared and used for H&E staining, IHC and IHF as described previously \textsuperscript{63, 113}. Lung sections used for estimating metastasis extent were prepared as we described previously \textsuperscript{77, 114}. For antibodies with a mouse origin, the M.O.M Mouse-on-Mouse Immunodetection kit (Vector Laboratories, Burlingame, CA) was used to block the immunostaining background from the endogenous mouse IgG. This study used the primary antibodies against Twist1 (ab50887), Foxa1 (ab23738, Abcam, Cambridge, MA), E-cadherin (610182, BD Bioscience, San Jose, CA), VIMENTIN (5741) and HA (3724, Cell Signaling, Danvers, MA). The secondary antibodies for IHC were biotinylated anti-Rabbit IgG and biotinylated anti-mouse IgG (Vector Laboratories, Burlingame, CA), which were used at a 1:400 dilution. The immunostaining signal was enhanced by using the VECTASTAIN ABC system (PK-6100, Vector Laboratories, Burlingame, CA), and then visualized by using the DAB kit (SK-4103, Vector Laboratories, Burlingame, CA). The tissue slides were counterstained with Harris Modified Hematoxylin and mounted with Permount. Double IHF was performed using the Tyramide Signal Amplification kit (Life technologies, Grand Island, NY) according
to the manufacturer's instructions. Stained tissue slides were examined and imaged under appropriate microscopes. Tumor and lung areas in the lung images were measured by using the NIH ImageJ software as described previously \(^7\), \(^{114}\).

### 2.7 Quantitative RT-PCR (qPCR)

Total RNA was isolated from WT/RCIP and Twist\(^{1TKO/RCIP}\) mammary tumors and BT549, Sum1315, MCF7, MCF7\(^{Ctrl}\), MCF7\(^{Twist1}\) and MCF7\(^{Twist1+Foxa1}\) cells using the TRIZOL reagent (Life technologies, Grand Island, NY). Reverse transcription was performed with 1 \(\mu\)g of RNA by using the Reverse Transcriptase Core kit (Eurogentec; Fremont, CA). qPCR was performed by using the matched Universal Taqman probes (Roche, Nutley, NJ) and gene-specific primer pairs (Table 2.1). The measurement of 18S or \(\beta\)-actin mRNA was used as an endogenous normalizer.

### 2.8 Genomic DNA extraction from paraffin embedded samples

Genomic DNA was isolated from lung paraffin sections of both groups of mice using QIAamp DNA FFPE Tissue Kit (56404, Qiagen, Hilden, Germany). The genomic DNA was used for PCR analysis of the Twist\(1\) gene knockout allele.
2.9 Micro-vascular density calculation

Microvascular density was determined as described previously. In brief, blood vessels were visualized by CD31 immunohistochemistry in the tissue sections prepared from mouse tumors. Tumor areas which contain greatest vessel density were defined as hot spots regions. Images of hot spots regions were taken under a microscope at 200× magnification. Any stained endothelial cells were counted to represent a single vessel if it was clearly separated from adjacent microvessels and other connective tissue elements.

2.10 Plasmids and cell transfection

For constructing the Foxa1-Luc promoter-reporter plasmid, the 1.5 kb 5’ regulatory sequence of the human Foxa1 gene was amplified by PCR using specific primers (Supplemental Table S1) from the genomic DNA of MCF7 cells and subcloned into the Kpn I and Nco I sites of the pGL3-basic-luciferase plasmid. The human E-cadherin-Luc and CSF1-Luc plasmids were described previously. The pCDNA-Twist1-2×flag expression plasmid was constructed by subcloning human Twist1 cDNA into the pCDNA-2×flag plasmid. The pCDH-Foxa1 plasmid was a gift from Dr. Bin He at Baylor College of Medicine.

For assaying the activity of a promoter-reporter construct, MCF7 cells were transfected
with the expression vectors and promoter-luciferase reporter plasmid as indicated in the Results section using the polyethylenimine reagent (23966-2, Polyscience, Niles, IL). The transfected cells were harvested 24 hours and lysed with the Reporter Lysis buffer (Promega, Madison, WI) for luciferase assay as described. The relative luciferase activity was normalized to the total amount of protein assayed.

To generate MCF7\textsuperscript{Twist1} cell lines, MCF7 cells were transfected with pCDNA-2×flag and pCDNA-Twist1-2×flag plasmids using the Polyethylenimine reagent. Transfected cells were cultured in DMEM medium containing 2 µg/ml of Hygromycin for 2 weeks. To generate MCF7\textsuperscript{Twist1+Foxa1} cell lines, two MCF7\textsuperscript{Twist1} cell lines were transfected with pCDH-Foxa1 plasmid and growth-selected for 14 days in DMEM medium containing 4 µg/ml of Puromycin. The surviving colonies in each experiment were individually isolated, expanded in growth medium and analyzed by Western blot.

In siRNA-based knocking down experiments, non-targeting control siRNAs and siRNAs targeting human Twist1 and c-Fos mRNAs were purchased from Dharmacon (Lafayette, CO). BT549, Sum1315 and MCF7 cells were transfected with siRNAs using the HiPerFect Transfection Reagent (301705, QIAGEN, Valencia, CA).

Knockdown of Twist1 mRNA by stable expression of shRNAs: pGIPZ-shCtrl lentiviral vector coding for a non-targeting shRNA (5’-CTTACTCTGCCCCAAGCGAGAG) and 2 pGIPZ-shTwist1 lentiviral vectors coding for two different Twist1 mRNA-targeting
sequences (TGAATGCATTAGACACCG and AGAGGAAGTCGATGTACCT) were obtained from the Cell-Based Assay Screening Service Core at Baylor College of Medicine. Lentivirus was packaged in 293T cells transfected with pMD2.G, psPAX2, and pGIPZ shRNA vectors in the ratio of 1:3:4. The culture medium was refreshed 6 hours after transfection. 24 hours after transfection, the medium containing viral particles was collected and polybrene was added to a concentration of 4 µg/ml for infecting cells. SUM1315 and MDA-MB-436 cells were infected with the prepared viral particles and selected in the culture medium containing 1 µg/ml of puromycin for 10 days. Survived cells were expanded and used for experimental analysis.

2.11 ChIP assay

The DNA-bound proteins in BT549, Sum1315, MCF7Ctrl and MCF7Twist1 cells were cross-linked using 1% formaldehyde for 10 minutes. ChIP assays were performed as described previously 68, 106. Antibodies used in these ChIP assays were for Twist1 (ab50887), c-JUN (ab31419, Abcam, Cambridge, MA), MTA2 (28791), HDAC2 (7899x, Santa Cruz, Dallas, TX), c-FOS (2250, Cell Signaling, Danvers, MA), Histone H3K9ac (39585, Active motif, Carlsbad, CA), RNA pol II (PLA0127, Sigma-Aldrich, St. Louis, MO) and M2 Flag antibody beads (A2220, Sigma-Aldrich, St. Louis, MO). ChIP-grade mouse IgG and rabbit IgG (Abcam, Cambridge, MA) were used as negative controls. The sequences of PCR primers used for amplifying the precipitated DNA samples are listed in Table 2.1.
2.12  **Cell migration and invasion assays**

Cell migration and invasion capabilities were measured as described previously. Briefly, individual cell migration was directly traced in a 96-well plate for 18 hours using the Cell Motility HCS Reagent kit (K08-000-11, Thermo Scientific, Rockford, IL). The track areas on the electronic images were quantitatively measured using the Image J software. Cell invasion was assayed by using BioCoat Matrigel Invasion Chambers (BD Biosciences, San Jose, CA). A total of 25,000 cells were seeded in each upper chamber with serum-free DMEM medium and the lower chamber was filled with DMEM containing 5% FBS. After cultured for 24 hours, non-migrating cells on the upper chambers were removed by a cotton swab, and cells invaded through the Matrigel layer to the underside of the membrane were stained with crystal violet and counted.

2.13  **In vivo metastasis assay**

MCF7Ctrl, MCF7Twist or MCF7Twist1+Foxa1 cells (1x10⁶) were injected into 8-week-old female SCID mice through the tail vein. These mice were sacrificed in 4 weeks after the injection. Their lung tissues were collected, photographed and processed for paraffin section. Metastasis was evaluated by counting the visible metastatic foci on the lung surface and measuring metastasis tumor area versus the lung area on H&E-stained lung sections as described previously. ⁷⁷, ¹¹⁴
2.14 Statistical analyses

All data were collected from several independent experiments. Each assay was performed in triplicate whenever applicable. All data were expressed as Mean ± SEM. Prism 4 Software (GraphPad, La Jolla, CA) was used to perform Student’s t test to analyze the data sets of mRNA level, tumor volume, luciferase activity, cell migration and invasion, metastatic tumor number and index. Log-rank test was used to analyze the data sets of mouse tumor-free survival and human breast cancer patient DMFS. Fisher’s exact test was used to analyze the data sets of Foxa1 IHC scores. Person’s Correlation test was used to analyze the Twist1 and Foxa1 expression data sets. In all statistical analyses, $p < 0.05$ was considered significant.
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**Table 2.1. Oligonucleotide primers and probes for PCR**
3. RESULTS

3.1 A new mouse model system for simultaneous deletion of the floxed alleles and expression of an oncogene in a subset of the mouse mammary epithelial cells

Du et al. has previously generated MMTV-TVA mouse line, in which the transgene for the \( tva \) avian virus subgroup A receptor is driven by MMTV (mouse mammary tumor virus) promoter and specifically expressed in the mammary gland luminal epithelial cells \(^{105}\). Intraductal instillation of the avian leukosis virus RCAS (replication-competent avian sarcoma-leukosis virus LTR splice acceptor) carrying PyMT caused the infection of a small subset of the TVA-expressing mammary epithelial cells in MMTV-TVA mice, resulting in rapid mammary tumorigenesis \(^{105}\). In order to inactivate a floxed gene of interest in the mammary tumor cells induced by the MMTV-TVA/RCAS-PyMT system, we constructed the RCAS-Cre-IRES-PyMT (RCIP) vector (Fig. 3.1A) and generated RCIP avian virus with this vector. Western blot analysis revealed that both Cre and PyMT proteins were expressed in the RCIP virus-infected DF1 cells but not in the RCAS control virus-infected DF1 cells (Fig. 3.1B). These results demonstrate that the RCAS promoter is strong enough to drive both Cre and PyMT protein expression after the viral DNA is integrated into the cell genome.

* Part of the data reported in this section is reprinted from “Twist1 Promotes Breast Cancer Invasion and Metastasis by Silencing Foxa1 Expression” by Yixiang Xu et al, 2016. Oncogene, Epub ahead of print. Yixiang Xu retains the copyright of the article as stated on the publisher’s website.
Next, we generated MMTV-TVA;ROSA26R bi-genic mice by crossbreeding MMTV-TVA and ROSA26R mice and injected RCIP avian virus into the mammary gland ducts of these mice. The ROSA26R locus contains a floxed transcriptional STOP sequence that silences the β-galactosidase (β-gal) expression and thus, serves as a reporter for Cre activity \(^{107,108}\). The mammary glands were isolated on the 3\(^{rd}\), 21\(^{st}\) and 48\(^{th}\) days post viral injection and analyzed by X-gal staining and histopathological examination (Fig. 3.1C). In the virus-injected mammary glands of MMTV-TVA;ROSA26R female mice, individually scattered β-gal-positive mammary gland epithelial cells were detected by day 3. One to multiple epithelial hyperplasia lesions consisting of only the β-gal-positive cells were developed by day 21 and invasive mammary gland tumors with β-gal-positive cells were formed by day 48 (Fig. 3.1D). Double immunofluorescent staining revealed that both PyMT and β-gal proteins were co-expressed in these tumor cells, indicating these tumor cells were derived from RCIP virus-infected mammary epithelial cells and Cre expressed in these cells worked effectively (Fig. 3.1E). In contrast, no β-gal-positive or PyMT-positive mammary epithelial cells or tumor cells were observed in the MMTV-TVA;ROSA26R mouse mammary glands without viral injection or in the ROSA26R mouse mammary glands with injection of RCIP virus (data not shown). These results demonstrate that the MMTV-TVA/RCIP (hereafter designated as TVA/RCIP) system is fully capable to express both Cre and an oncogene in the same mammary epithelial cells for simultaneous and effective deletion of a floxed gene and induction of tumorigenesis.
Figure 3.1. The mammary tumorigenesis mouse model induced by the novel MMTV-TVA/RCIP system. A. The RCAS-Cre-IRES-PyMT-HA (RCIP) vector of the avian virus. B. Detection of both Cre and PyMT-HA proteins in the RCIP vector-transfected DF1 cells by Western blotting, while these proteins were not present in the untransfected DF1 (Ctrl) cells. C. The strategies to induce β-gal-labeled mammary tumors in the female MMTV-TVA;ROSA26R mice by using the RCIP system. D. X-gal staining to show RCIP virus-infected and induced mammary tumor cells (blue color) in the mammary glands of MMTV-TVA;ROSA26R mice after intraductal injection of RCIP virus for 3, 21 and 48 days. E. Double immunofluorescence (IF) staining for PyMT-HA (red) and β-gal (green) in a RCIP-induced mammary tumor isolated from MMTV-TVA;ROSA26R mice. A section of a mammary gland isolated from a normal female mouse served as a negative control for HA and β-gal IF staining.
3.2 Tumor specific knock-out of Twist1 using TVA/RCIP system in Twist1-floxed mice

To knockout Twist1 in PyMT-induced mammary tumor cells by using TVA/RCIP system, we generated female MMTV-TVA and MMTV-TVA; Twist1\(^{F/F}\) mice with the same strain background by crossbreeding MMTV-TVA and the floxed Twist1 (Twist1\(^{F/F}\)) mice \(^{63, 105, 109}\). To get Twist1\(^{TKO}\)/RCIP tumors (RCIP-induced tumors with tumor cell-specific Twist1 knockout), we injected RCIP virus into the ductal lumens of either one or two of their 4\(^{th}\) mammary glands, when MMTV-TVA; Twist1\(^{F/F}\) mice were 8 weeks old. PyMT oncogene induces tumorigenesis only in the Twist1 knockout mammary epithelial cells (Fig. 3.2A). MMTV-TVA mice were used as control group to get WT/RCIP tumors.

To evaluate Twist1 knockout efficiency, we prepared genomic DNA, RNA and protein samples from RCIP virus-induced mammary tumors, and performed genotype, Q-PCR and Western blot analyses. As expected, genotype analysis only detected wild type Twist1 allele in WT/RCIP tumors, but detected both the deleted and the floxed Twist1 alleles in Twist1\(^{TKO}\) tumors due to the presence of both the tumor cells with Twist1 deletion and the non-tumor cells with the floxed Twist1 in the tumor tissues (Fig. 3.2B). Q-PCR analysis detected a 14-fold decrease in Twist1 mRNA expression in Twist1\(^{TKO}\)/RCIP tumors compared with WT/RCIP tumors (Fig. 3.2C). Western blot analysis revealed a significant reduction of Twist1 protein in Twist1\(^{TKO}\)/RCIP tumors.
versus WT/RCIP tumors (Fig. 3.2D). We further performed immunostaining and confirmed that variable levels of Twist1 protein was expressed in a sub-population (~6%) of tumor cells and certain stromal cells in WT/RCIP tumor tissues, but Twist1 protein was only detected in the stromal cells in $Twist1^{TKO}$/RCIP tumor tissues. Moreover, Twist1 and PyMT-HA double positive tumor cells were observed in WT/RCIP tumors, while only PyMT-HA single positive tumor cells were seen in $Twist1^{TKO}$/RCIP tumors (Fig. 3.2E). These results demonstrate that Twist1 is expressed in a subset of mammary tumor cells and certain stromal cells in WT/RCIP tumors, while in $Twist1^{TKO}$/RCIP tumors Twist1 was completely knocked out in the tumor cells but normally expressed in the infiltrating stromal cells.
Figure 3.2. Mammary tumor cell-specific knockout of Twist1 in mice. A. Experimental strategies to induce mammary tumors with and without TKO of Twist1 using the MMTV-TVA/RCIP system. TVA, MMTV-TVA mice; WT, wild type mice. B. PCR-based analysis of the wild type (WT), floxed (Twist1^flox^) and Cre-deleted (Twist1^TKO^) Twist1 alleles in WT/RCIP and Twist1^TKO^/RCIP mouse mammary tumors. C. Q-PCR-based analysis of Twist1 mRNA in WT/RCIP and Twist1^TKO^/RCIP mouse mammary tumors. Data are presented as Mean ± SEM. D. Western blot analysis of Twist1 protein in WT/RCIP (n=4) and Twist1^TKO^/RCIP (n=4) mammary tumors. The β-actin served as an endogenous loading control. The mouse IgG band was recognized by the secondary antibody. E. Detection of Twist1 (brown) by IHC in both tumor and stromal cells in WT/RCIP tumors but only in stromal cells in Twist1^TKO^/RCIP tumors (upper panels). In lower panels, double IF staining detected Twist1 (arrowheads) in PyMT-HA-expressing tumor cells in WT/RCIP tumors, but no Twist1 in PyMT-HA-expressing tumor cells in Twist1^TKO^/RCIP tumors. Scale bars, 50 µm.
3.3 Specific ablation of Twist1 in the PyMT-induced mammary tumor cells does not affect primary tumor development in mice

Palpable mammary tumors were detected in both genotype groups of mice within 6-10 weeks after viral injection, showing no difference in the latency of tumor initiation between MMTV-TVA;\textit{Twist1}^{F/F} and MMTV-TVA mice (Fig. 3.3 A). Thirty-five and 26 tumors were developed from 60 virus-infected mammary glands in MMTV-TVA;\textit{Twist1}^{F/F} mice and 42 virus-infected mammary glands in MMTV-TVA mice, respectively. The tumorigenic rate was about 60\% for both groups. These results indicate that specific inactivation of \textit{Twist1} in the mammary epithelial cells does not affect PyMT-induced mammary tumor formation.

We measured tumor growth for 10 weeks after a palpable tumor was detected in the RCIP virus-infected mammary gland of MMTV-TVA;\textit{Twist1}^{F/F} or MMTV-TVA mice. The tumor growth data was only collected from mice bearing a single tumor. We found that the average tumor growth rates in the two groups of mice were very similar (Fig. 3.3B). Cell proliferation rates assayed by Ki67 immunohistochemistry also showed no significant difference between \textit{Twist1}^{TKO}/RCIP mammary tumors isolated from MMTV-TVA;\textit{Twist1}^{F/F} mice and WT/RCIP mammary tumors (RCIP-induced tumors with wild type \textit{Twist1}) isolated from MMTV-TVA mice (Fig. 3.4A). Analysis of cell apoptosis by cleaved-caspase 3 staining and macrophage recruitment by F4-80 staining also revealed no difference between these two groups (Fig. 3.4 B&C). These results indicate
that the autonomous cellular function of Twist1 is not required for the growth of PyMT-induced mammary tumors in mice.

Figure 3.3. Tumor cell-specific knockout (TKO) of Twist1 does not influence tumor initiation or growth in mice. A. Tumor-free survival curves for WT/RCIP (RCIP-infected MMTV-TVA mice) and Twist1TKO/RCIP (RCIP-infected MMTV-TVA;Twist1F/F mice) mice. Mammary tumor development was monitored by palpation. B. WT/RCIP and Twist1TKO/RCIP mammary tumor growth curves generated by measuring tumor volumes (Vol.) once a week after palpable tumors were detected.

3.4 Specific ablation of Twist1 in the PyMT-induced mammary tumor cells inhibits lung metastasis

Next, we examined lung metastasis in the RCIP virus-infected MMTV-TVA and MMTV-TVA;Twist1F/F mice in which a palpable tumor had grown for 10 weeks. We observed lung metastases in 67% (12/18) of MMTV-TVA mice with 5.38 metastatic foci/lung in average number. However, we only observed lung metastases in 39% (7/18) of MMTV-TVA;Twist1F/F mice with 0.38 metastatic foci/lung in average (Fig. 3.5A). On the lung tissue sections, the
Figure 3.4 IHC staining of primary tumors. A. Representative images of Ki67 IHC on WT/RCIP and Twist1TKO/RCIP primary tumor sections and the quantification of staining result. B. Representative images of c-Caspase3 (cleaved caspase3) IHC on WT/RCIP and Twist1TKO/RCIP primary tumor sections and the quantification of staining result. C. Representative images of F4-80 IHC on WT/RCIP and Twist1TKO/RCIP primary tumor sections and the quantification of staining result.
average percentage of metastasis area to total lung area, which is an index for metastasis extent, was also drastically reduced in MMTV-TVA;\textit{Twist1}\textsuperscript{F/F} mice versus MMTV-TVA mice (Fig. 3.5B). These results demonstrate that \textit{Twist1} null mammary tumor cells have a drastically decreased metastasis capability compared with \textit{Twist1} WT mammary tumor cells.

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\caption{\textbf{Figure 3.5. Tumor cell-specific knockout (TKO) of \textit{Twist1} inhibits lung metastasis in mice.} \textbf{A.} The images of lungs isolated from MMTV-TVA/RCIP and MMTV-TVA;\textit{Twist1}\textsuperscript{F/F}/RCIP mice bearing palpable mammary tumors for 10 weeks (upper panels). Arrowheads indicate metastasis foci observed on the lung surface. The numbers of metastasis foci observed in the sections of individual lungs of both mouse groups are presented in the lower panel. \textbf{B.} H&E-stained lung sections of mice with WT/RCIP and \textit{Twist1}\textsuperscript{TKO}/RCIP mammary tumors for 10 weeks. Arrowheads indicate metastasis foci (upper panels). Metastasis index was presented as the average percentage of metastasis tumor areas to total lung areas measured on the electronic images recorded from H&E-stained lung sections. Scale bars, 100 µm.}
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We also performed Ki67 immunostaining and found higher proliferating ratio in MMTV-TVA;\(Twist1^{E/F}\) mice metastasis tumors than in MMTV-TVA mice metastasis tumors (Fig. 3.6A). This result suggests that the reduced metastasis of Twist1 null tumor cells in MMTV-TVA;\(Twist1^{E/F}\) mice was not caused by any decrease in tumor cell proliferation in the lung. Instead, the lung metastasis may happen at a later stage in MMTV-TVA;\(Twist1^{E/F}\) mice versus MMTV-TVA mice.

We isolated genomic DNA from lung paraffin section with metastasis foci. Genotype analysis showed wild type \(Twist1\) allele is only detected in WT/RCIP groups and floxed allele is only detected in \(Twist1^{TKO}\) group (Fig. 3.6B). However, deleted \(Twist1\) alleles were detected in some \(Twist1^{TKO}\) lungs, suggesting that \(Twist1\) null tumor cells could also metastasize to lung but have a drastically decreased metastasis capability compared with \(Twist1\) WT mammary tumor cells.
3.5 Specific ablation of Twist1 in the PyMT-induced mammary tumor cells inhibits EMT and invasiveness

Cell line based studies showed that Twist1 promotes EMT, angiogenesis and local invasiveness. Thus, we examined EMT status of both Twist1TKO/RCIP and WT/RCIP tumors by triple fluorescent IHC staining of Twist1/PyMT-HA/E-cadherin or Twist1/PyMT-HA/Vimentin. We found that Twist1 positive tumor cells (Twist1/PyMT-HA double positive) showed increased Vimentin expression and decreased E-cadherin...
expression compared with Twist1 negative tumor cells in WT/RCIP tumor sections (Fig. 3.7A and B). On the other hand, we did not see any Twist1 positive tumor cells from the Twist1TKO/RCIP tumor sections. However, we did see a few Twist1 negative tumor cells from both groups showed decreased E-cadherin or increased Vimentin (Fig. 3.7 A and B), suggesting that Twist1 is not the only regulator of EMT in the tumor cells.

Quantification of the staining showed there are about 9.0 % Vimentin positive tumor cells and about 45% of which are Twist1 positive in WT group. In Twist1TKO/RCIP tumors, about 2.6% of tumor cells are Vimentin positive. Vimentin and E-cadherin expression levels in Twist1 positive and negative tumor cells in WT/RCIP group were scored in a scale of 0-3 (0 means negative/non-detectable while 3 means strong expression). Twist1 positive tumor cells showed significant higher Vimentin score but lower E-cadherin score. These data suggest that Twist1 is a master inducer of EMT in the mouse mammary gland tumors.
Figure 3.7. Vimentin and E-cadherin expression in Twist1 positive and negative tumor cells. A. IF staining of Vimentin, PyMT-HA and Twist1 in WT/RCIP and Twist1TKO/RCIP mammary tumors. Vimentin expression level in Twist1 positive tumor cells and Twist1 negative tumor cells were scored and shown in right panel. B. IF staining of E-cadherin, PyMT-HA and Twist1 in WT/RCIP and Twist1TKO/RCIP mammary tumors. E-cadherin expression level in Twist1 positive tumor cells and Twist1 negative tumor cells were scored and shown in right panel. Scale bar, 50 µm.

Micro-vascular density is an important marker for invasiveness, thus we performed CD31 IHC on both Twist1TKO/RCIP and WT/RCIP tumor sections. Knockout of Twist1 in tumor cells significantly decreased Micro-vascular density by around 50% (Fig. 3.8A). Consistent with the active EMT and angiogenesis promoted by Twist1, we detected more circulating tumor cells (CTCs) from the blood of mice with WT/RCIP tumors compared with those mice with Twist1TKO/RCIP tumors (Fig. 3.8B). Taken
together, Twist1 promotes EMT and angiogenesis to facilitate primary mammary gland tumor cell invasiveness and intravasation into the blood circulation.

Figure 3.8. TKO of Twist1 in the PyMT-induced mammary tumor cells inhibits angiogenesis and invasion into blood circulation. A. Micro-vascular density of WT/RCIP and Twist1TKO/RCIP mouse mammary tumors. B. Circulating tumor cells (CTCs) colony culture from mice with WT/RCIP or Twist1TKO/RCIP mammary tumors.

3.6 Twist1 expression is turned off in lung metastasis foci

To figure out Twist1’s role in metastasis tumors, we examined Twist1 expression in lung metastasis foci. Interestingly, double staining of Twist1 and tumor marker PyMT-HA showed that metastasis foci do not express Twist1 protein in either WT/RCIP or Twist1TKO/RCIP groups (Fig 3.9A). This phenotype is consistent with the previous report that Twist1 is turned off in metastasis tumors through MET49. We then performed double staining of PyMT-HA with E-cadherin or Vimentin and found that all metastasis
tumor cells are E-cadherin positive but Vimentin negative. Double staining of E-cadherin and Vimentin also showed no overlapping, further supports the conclusion that Twist1 is turned off in the metastasis foci and metastasis tumor showed epithelial phenotype. (Fig. 3.9B, C and D).

3.7 Twist1 is expressed at late tumor stage and is associated with reduced ER and Foxa1 expression in mouse tumors

Above data showed that Twist1 was non-essential in early stage of tumor progression, however, it was unclear whether Twist1 is expressed but does not contribute to early stage tumor progression or it is not expressed at all. Thus we examined Twist1 protein expression by IHC in both groups of tumors at early stage (6 weeks after tumor became palpable, volume ≈ 0.5 cm³) and late stage (10 weeks after tumor became palpable, volume ≈ 1.0 cm³). Interestingly, we found no Twist1 immunoreactivity was detected in the early stage tumor cells, while ERα protein was detected at high levels in both tumor cells and non-tumor mammary epithelial cells (Fig. 3.10A). In late stage tumors, about 6% of WT/RCIP tumor cells express Twist1, suggesting that Twist1 mainly functions at late stage.
Figure 3.9. Twist1, Vimentin and E-cadherin expression in lung metastasis tumor foci of WT/RCIP and Twist1TKO/RCIP groups. A. IF staining of Twist1 and PyMT-HA in lung metastasis foci of both groups. B. IF staining of E-cadherin and PyMT-HA in lung metastasis foci of both groups. C. IF staining of Vimentin and PyMT-HA in lung metastasis foci of both groups. D. IF staining of E-cadherin and Vimentin in lung metastasis foci of both groups.
Figure 3.10. Twist1, ERα and Foxa1 IHC in mouse tumors and IRS score. A. IHC for Twist1 ERα and Foxa1 (brown) in WT/RCIP and Twist1TKO/RCIP mammary tumors at 6 and 10 weeks after becoming palpable in mice. B. The sample distribution and Fisher’s Exact test of ERα (strong positive (IRS>4) and weak positive or negative (IRS ≤ 4)) and Foxa1 (negative (IRS=0) and positive (IRS>0)) in WT/RCIP and Twist1TKO/RCIP mouse mammary tumors at 10 weeks after becoming palpable in mice.

As a semi-quantitative analysis, we evaluated the immunoreactivity scores (IRS) of ERα and Foxa1 in both groups of tumors using the Allred Scoring System. In late stage tumors ERα immunoreactivity was drastically decreased compared with that detected in the early stage (20% vs 58% positivity). Interestingly, although ERα was also reduced in
most of the tumor cells of late stage $Twist1^{TKO}$/RCIP tumors compared with early stage, there were still 59% of tumor cells expressing detectable ER$\alpha$ and 14% of tumor cells expressing high level ER$\alpha$. As an ER$\alpha$ co-activator and a determinant of ER$\alpha$ signaling in breast cancer, Foxa1 protein showed similar expression pattern as ER$\alpha$. Foxa1 was detected at high levels in both tumor cells and non-tumor mammary epithelial cells at early stage (Fig. 3.10A). However, Foxa1 immunoreactivity was very weak (12%) or negative (71%) in WT/RCIP tumor cells, while there were still 71% of tumor cells with detectable Foxa1 and 8% of tumor cells with high level Foxa1 in large $Twist1^{TKO}$/RCIP tumor group. Using Allred Scoring System, we found that late stage tumors are still ER$\alpha$ positive (IRS > 2, as generally accepted for determining positivity of ER$\alpha$ in human breast cancer patients), thus we compared ER$\alpha$ strong positive (IRS>4) to ER$\alpha$ weak positive or negative (IRS $\leq$ 4) ratios in the two groups of tumors and found that $Twist1^{TKO}$/RCIP tumors are all ER$\alpha$ strong positive while more than 50% of WT/RCIP tumors turn to ER$\alpha$ weak positive or negative (Fig. 3.10B left panel). On the other hand, Foxa1 expression level is low in both groups of tumors, thus we compared Foxa1 negative (IRS = 0) to Foxa1 weak or strong positive (IRS > 0) ratios between the two groups. As shown in the right panel of Fig. 3.10B, the ratio of Foxa1-positive tumors was significantly higher in $Twist1^{TKO}$/RCIP tumor group versus WT/RCIP tumor group. Together, these results demonstrate that the level of Twist1 expression negatively correlates with the level of ER$\alpha$ and Foxa1 expression in the PyMT-induced mouse mammary tumors. ER$\alpha$ has been shown to be a direct target of Twist1 in human breast cancer cells, thus we next investigated whether and how Foxa1 is regulated by Twist1.
3.8 Twist1 expression inversely associates with Foxa1 expression in human breast cancer

Based on the mouse tumor IHC result, we hypothesized that Twist1 inhibits Foxa1 expression to promote breast cancer invasiveness. To identify whether this regulation is true in human breast cancer, we analyzed a dataset deposited in the NCBI GEO database (GSE53222) and found that Twist1 expressed in the T47D ER-positive breast cancer cells decreased the expression of a group of luminal epithelial genes including Foxa1, ESR1, PRLR, PGR, MUC1 and CLDN3 (Fig. 3.11), further supporting the hypothesis that Foxa1 is target of Twist1 in breast cancer.

![Figure 3.11. Downregulated genes by Twist1 overexpression in T47D human breast cancer cells.](image)

Several known luminal type breast cancer marker genes are downregulated by Twist1 overexpression, including Foxa1, ESR1, GATA3, MUC1. (Data source: Shi J et al, Cancer Cell, 2014)

We further analyzed the correlation between the expression levels of Twist1 and Foxa1
mRNAs in human breast cancer using another dataset from 281 human breast tumors in NCBI GEO (GSE2034)\textsuperscript{118}. This analysis also revealed a significant negative correlation between the levels of \emph{Twist1} and \emph{Foxa1} mRNA expression (Fig. 3.12A). We also analyzed the correlation between the expression levels of \emph{Twist1} and \emph{Foxa1} mRNAs in luminal and basal subtype human breast cancer cell lines by using GEO2R software and a dataset in the NCBI GEO database (GSE15361)\textsuperscript{119}. Among 44 breast cancer cell lines with RNA-profiling data, the level of \emph{Twist1} mRNA negatively correlated with the level of \emph{Foxa1} mRNA (Fig. 3.12D). The average level of \emph{Twist1} mRNA was significantly higher in basal-like breast cancer cell lines compared with luminal type of breast cancer cell lines (Fig. 3.12B). In contrast, the average level of \emph{Foxa1} mRNA was significantly lower in basal-like breast cancer cell lines compared with luminal type of breast cancer cell lines (Fig. 3.12C). To validate these data at protein levels, we performed Western Blot for Twist1 and Foxa1 in a group of breast cancer cells lines. We found that Twist1 is highly expressed in basal-like breast cancer cell lines, while Foxa1 is exclusively expressed in luminal type breast cancer cell lines. We further analyzed a cohort of 276 human breast tumors by IHC staining of both Twist1 and Foxa1. Twist1 protein was detected in 13 out 276 (5\%) tumors and Foxa1 is detected in 94\% tumors (245 out of 261), indicating that this cohort mainly consists of luminal breast tumors. Interestingly, 179 out of 261 (68.6\%) tumors showed very high (>6) Foxa1 immunoreactive scores (IRSs). However, the 13 Twist1-positive tumors had IRSs of 6 or lower. The immunoreactive scores of Twist1 and Foxa1 and the ER\(\alpha\), PR and HER2 expression profiles for these 13 Twist1 positive tumors are provided in Table 3.1. Among these 13
Twist1-positive tumors, 5 tumors showed significantly decreased Foxa1 protein compared with Twist1-negative tumors (Fig. 3.12F). Twist1 protein levels were also negatively correlated with Foxa1 protein levels among these 13 Twist1-positive tumors (Fig. 3.12G). These results demonstrate that the levels of Twist1 expression are negatively correlated with the levels of Foxa1 expression in human breast tumors. Thus we proposed that Twist1 could directly silence Foxa1 expression to promote de-differentiation and metastasis of breast cancer cells.

**Figure 3.12.** *Twist1* expression is negatively associated with *Foxa1* expression in human breast cancers. A. Pearson’s Correlation test of *Twist1* and *Foxa1* mRNA expression levels in a published data set of human breast tumor cohort. B and C. Expression levels of *Twist1* (Panel B) and *Foxa1* (Panel C) mRNAs in luminal and basal breast cancer cell lines. Data analysis was based on a published data set. D. Pearson’s Correlation analysis between *Twist1* and *Foxa1* mRNA expression in breast cancer cell lines shown in Panels B and C. E. Western blotting analysis of Twist1 and Foxa1 in human breast cancer cell lines. F and G. Representative images of IHC for Twist1 and Foxa1 in human breast tumors. Adjacent tumor sections were used for Twist1 and Foxa1 IHC (Panel F). Twist1 and Foxa1 protein levels are inversely correlated in Twist1-positive breast tumors. The r and p values were obtained by Pearson’s correlation analysis (Panel G). The scale bar in Panel F, 50 µm.
Table 3.1 Clinical information for Twist1-positive breast tumors. ER, estrogen receptor; PR, progesterone receptor; HER2, human epithelial growth factor receptor 2; IRS, immunoreactivity score; N, negative; P, positive

3.9 Twist1 directly silences Foxa1 mRNA transcription by recruiting NuRD complex

The luminal type breast cancer cell line MCF7 expresses ERα and Foxa1, but not Twist1. To establish a cellular model for investigating the molecular mechanisms responsible for Twist1 to regulate Foxa1 expression, we generated stable MCF7 cell lines with either the empty control vector (MCF7Ctrl) or the Twist1-expressing vector (MCF7Twist1). Twist1 expression in MCF7Twist1 cells drastically decreased Foxa1 mRNA and protein levels (Fig. 3.13A). Conversely, knockdown of Twist1 mRNA in the basal-
like SUM1315 human breast cancer cells with high Twist1 and low Foxa1 expressions moderately increased the expression levels of Foxa1 mRNA and protein (Fig. 3.13B). However, ectopic expression of Foxa1 in either SUM1315 or BT549 breast cancer cells did not alter the expression levels of Twist1 mRNA and protein (Fig. 3.13 C and D). These results indicate that Twist1 suppresses Foxa1 expression, while Foxa1 does not regulate Twist1 expression.

Figure 3.13. Twist1 silences Foxa1 expression in breast cancer cells. A. Expression of Twist1 in MCF7 cells reduced Foxa1 mRNA and protein as measured by Q-PCR and Western blotting. B. Knockdown of Twist1 mRNA in SUM1315 cells by using Twist1 siRNA (siTwist1) increased Foxa1 mRNA and protein. C and D. Overexpression of Foxa1 did not influence Twist1 mRNA and protein in either BT549 or SUM1315 cells. All experiments were repeated 3 times and the representative data are presented. *, P<0.05 by Student’s t test.

To examine whether Twist1 was recruited to the enhancer/promoter regions of the
Foxa1 gene, we performed chromatin immunoprecipitation (ChIP) assays by using a Twist1 antibody to pull down Twist1-associated genomic DNA in BT549 breast cancer cells and PCR to measure specific DNA regions of the Foxa1 enhancer or promoter. We found that Twist1 was associated with the 5’ regulatory region at -1 kb from the transcriptional starting site (TSS), but it was not associated with -2, -3, -4 and -5 kb regions (Fig. 3.14A). To examine the specific role of Twist1 in regulating the transcriptional activity of Foxa1 promoter, we constructed a Foxa1 promoter-luciferase (Foxa1-Luc) reporter containing 1.46-kb 5’ regulatory sequence with the Twist1-binding region (Fig. 3.14B). Expression of Twist1 significantly decreased the activity of Foxa1-Luc reporter in MCF7 cells. As expected for positive and negative controls, Twist1 decreased the activity of CDH1-Luc reporter, but did not affect the activity of CSF1-Luc reporter (Fig. 3.14C). These results indicate that Twist1 is recruited to the 5’ regulatory region proximal to the Foxa1 promoter.

In silico analysis of the proximal Foxa1 promoter region revealed 11 E-boxes. Twist1 is more likely to bind two (EB3 and EB7) of these E-boxes as predicted by MatInspector Software (Genomatix) (Fig. 3.14B). Deletion of either EB3 site in Foxa1-Luc (EB3M-Luc) reporter or EB7 site in Foxa1-Luc (EB7M-Luc) reporter significantly relieved Twist1-repressed Foxa1 promoter activity in MCF7 cells. Double deletions of both EB3 and EB7 sites in Foxa1-Luc (EB3/7M-Luc) reporter additively relieved Twist1-repressed Foxa1 promoter activity (Fig. 3.14B and D). Since the activity of EB3/7M-Luc reporter was still partially silenced by Twist1 in MCF7 cells, Twist1 might also bind to other E-
Figure 3.14. Twist1 directly bind to E-box element of Foxa1 promoter. A. ChIP assays performed with BT549 breast cancer cells and Twist1 antibody. Non-immune IgG served as a negative control. Immunoprecipitated DNA was assayed by real time PCR with primers specific to the Foxa1 5’ regulatory sequences at the indicated locations. B. The wild type and mutated constructs of the Foxa1 promoter-luciferase reporter (Foxa1-Luc). EB3M-Luc, EB7M-Luc and ER3/7M-Luc had mutated E-box3, E-box7 or both of them, respectively. C & D. Cell transfection assays. Twist1 expression in MCF7 cells repressed the activities of Foxa1-Luc (the tester) and CDH1-Luc (a positive control), while it did not affect the activity of CSF1-Luc (a negative control) (Panel C). Twist1 showed much less repression on the activities of EB3M-Luc, EB7M-Luc and EM3/7M-Luc versus Foxa1-Luc in MCF7 cells (Panel D). Control cells were transfected with equal amount of the empty vector DNA. *, P< 0.05 by Student’s t test.
boxes to repress the Foxa1 promoter. On the other hand, in SUM1315 cells with endogenous Twist1 expression, mutation of EB3 and/or EB7 increased Foxa1-Luc reporter activity, while knockdown of Twist1 resulted in lesser fold increases in the mutant Foxa1 promoter activities versus the wild type promoter (Fig. 3.15 A and B). These results demonstrate that Twist1 can use both EB3 and EB7 sites to silence the transcriptional activity of the Foxa1 promoter. We previously reported that Twist1 recruits nucleosome remodeling deacetylase (NuRD) complex to repress CDH1 and estrogen responsive program (E-Reps). These results suggest that Twist1 may use multiple E-boxes to silence transcriptional activity of Foxa1 promoter.

Figure 3.15. Mutation of Twist1-binding sites or knockdown of Twist1 mRNA increases Foxa1 promoter-luciferase (Luc) reporter activity in SUM1315 cells. A. Knockdown of Twist1 mRNA in SUM1315 cells increased the activities of Foxa1-Luc (the tester) and CDH1-Luc (a positive control), but did not affect the activity of CSF1-Luc (a negative control). B. SUM1315 cells with normal Twist1 (shCtrl) or with Twist1 knockdown (shTwist1) were transfected with wild type Foxa1 or mutant Foxa1 promoter-Luc reporters as indicated. Mutation of either E-box 3 (EB3M-Luc), E-box 7 (EB7M-Luc) or both of them (EB3/7M-Luc) in the promoter sequence relieved Twist1-mediated suppression and thus, increased the basal activities of the mutant promoters in control cells. For the same reason, knockdown of Twist1 expression showed less fold relieves of the three mutant Foxa1 promoter-Luc reporters when compared with the wild type Foxa1 promoter-Luc reporter. All experiments were repeated 3 times and the representative data with 3 technical replicates are presented as Mean ± SEM. *, P < 0.05 by Student’s t test.
receptor (ERα) promoters\textsuperscript{68, 120}. Here, our ChIP assays revealed that Twist1 was specifically recruited to the proximal promoter regions of both CDH1 and Foxa1 genes in MCF7\textsuperscript{Twist1} cells (Fig. 3.16A). The endogenous HDAC2 and MTA2 proteins, two of the NuRD complex components, were also recruited to the same chromatin regions of both CDH1 and Foxa1 genes in a Twist1 expression-dependent manner (Fig. 3.16 B and C). In agreement with the Twist1-recruited NuRD complex containing histone deacetylase activity, the levels of acetylated histone H3K9 (H3K9-ace) were markedly reduced in the same Twist1-binding regions of both CDH1 and Foxa1 genes in MCF7\textsuperscript{Twist1} cells versus MCF7\textsuperscript{Ctrl} cells (Fig. 3.16D). Accordingly, RNA polymerase II recruited to these gene promoters was significantly reduced in MCF7\textsuperscript{Twist1} cells versus MCF7\textsuperscript{Ctrl} cells (Fig. 3.16F). Furthermore, knockdown of Twist1 in SUM1315 cells decreased HDAC2 and MTA2 recruitments and increased H3K9-ace and RNA polymerase II recruitment on Foxa1 promoter (Fig. 3.17). These results demonstrate that Twist1 largely silences the transcriptional activity of the Foxa1 promoter by recruiting NuRD gene-repressing complex, the same mechanism as it represses CDH1 transcription.
Figure 3.16. Overexpressed Twist1 in MCF7 cells recruits NuRD complex to Foxa1 promoter to repress its expression. ChIP assays were performed in MCF7Ctrl and MCF7Twist1 stable cell lines using antibodies against Twist1 (Panel A), HDAC2 (Panel B), MTA2 (Panel C), H3K9-ace (Panel D) and RNA-P1I (Panel E). Non-immune IgG served as a negative control. Immunoprecipitated DNA was measured by real time PCR using primers for amplifying the Twist1-binding regions in the CDH1 gene (a positive control) and the Foxa1 gene (E-box3) as indicated. All experiments were repeated 3 times and the representative data are presented as Mean ± SEM.
Figure 3.17. Knockdown of Twist1 mRNA reduced MTA2 and HDAC2 recruitments but increased H3K9-Ace and RNA polymerase II (RNA-PII) recruitment to the Foxa1 promoter in SUM1315 cells. ChIP assays were performed with SUM1315 cells with normal Twist1 (shCtrl) or Twist1 knockdown (shTwist1) and with non-immune IgG (negative control) or antibodies against Twist1, HDAC2, MTA2, H3K9-ace or RNA-PII as indicated. The input and precipitated DNA samples were analyzed by real-time PCR that amplifies the E-box 3 region of the Foxa1 promoter. All experiments were repeated 3 times and the representative data with 3 technical replicates are presented as Mean ± SEM. *, $P < 0.05$ by Student’s $t$ test.

3.10 Twist1 silences Foxa1 expression through inhibiting AP-1-promoted activation of the Foxa1 promoter

Computational analysis of the Foxa1 promoter region identified two AP-1 binding sites conserved in both human and mouse Foxa1 genes (Fig. 3.18A). To define the role of AP-1 in regulating Foxa1 promoter, we expressed c-Jun and c-Fos and assayed their effects on the Foxa1-Luc reporter in MCF7 cells. Expression of c-Jun and c-Fos significantly increased the activity of Foxa1-Luc reporter in a dose-dependent manner (Fig. 3.18B). In contrast, knockdown of c-Fos in MCF7 cells drastically reduced the...
expression levels of Foxa1 mRNA and protein (Fig. 3.18C). Furthermore, deletion of either AP-1 site at bp -895 or bp -798 or both diminished AP-1-promoted activities of the *Foxa1* AP1M1-Luc, AP1M2-Luc and AP1M1/2-Luc reporters (Fig. 3.18D). These results demonstrate that AP-1 is a transcriptional activator of the *Foxa1* promoter.

We further performed ChIP assays to examine whether Twist1 could inhibit c-Fos from being recruited to the AP-1 sites of the *Foxa1* proximal promoter region. Indeed, Twist1 expression in MCF7 cells abolished the recruitment of c-Fos to the *Foxa1* proximal promoter region (Fig. 3.18E). As a control, Twist1 did not inhibit c-Fos recruitment to the *CSF1* promoter region.106 These results suggest that Twist1 plays a crucial role to repress AP-1-mediated activation of the *Foxa1* promoter by preventing AP-1 recruitment to the *Foxa1* proximal promoter region. As expected, expression of Twist1 in MCF7 cells markedly repressed the transcriptional activity of the Foxa1-Luc reporter. However, Twist1 expression did not have an obvious effect on the activity of the *Foxa1* promoter with deleted AP-1 binding sites, including AP1M1-Luc, AP1M2-Luc and AP1M1/2-Luc reporters (Fig. 3.18F). Consistent results were also observed in SUM1315 cells with *Twist1* knockdown (Fig. 3.19 A and B). These results suggest that Twist1 plays a role in silencing AP-1-mediated activation of the *Foxa1* promoter.
Figure 3.18. Twist1 inhibits AP1-mediated activation of the Foxa1 promoter by inhibiting AP1 recruitment.  A. The two conserved AP1 sites in both human and mouse Foxa1 promoter regions.  B. Luciferase activity of MCF7 cells transfected with Foxa1-Luc plasmid with the empty vector (-) or c-Jun and c-Fos expression vectors.  C. Q-PCR and Western blot assays showing Foxa1 mRNA and protein changes upon c-fos knockdown in MCF7 cells.  D. Luciferase assay of MCF7 cells co-transfected with the indicated luciferase reporter and the empty vector (-) or c-Jun and c-Fos expression vectors.  E. ChIP assays performed with MCF7^{Ctrl} and MCF7^{Twist1} cells and non-immune IgG (negative control) or c-Fos antibody as indicated.  The DNA fragment between the two arrowheads in Panel A was amplified by PCR from the immunoprecipitated DNA. Amplification of the known AP1-binding region by PCR in the CSF1 promoter served as a positive control.  F. Luciferase assay of MCF7 cells co-transfected with the indicated luciferase reporter and the empty vector (Ctrl) or Twist1 expression plasmid. Experiments were repeated 3 times and the representative data are presented as Mean ± SEM.  * in Panels B–D, $P<0.05$ by Student’s $t$ test.
Figure 3.19. Twist1 inhibits AP1-activated Foxa1 promoter activity in SUM1315 cells. A. Knockdown of Twist1 mRNA by shTwist1 increased c-Fos recruitment to the Foxa1 promoter. ChIP assays were performed with SUM1315 cells expressing non-targeting shRNA (shCtrl) or shTwist1 that targets Twist1 mRNA and with control IgG or c-Fos antibody as indicated. B. Mutation of AP-binding sites in the Foxa1 promoter decreased the promoter activity and knockdown of Twist1 expression markedly increased wild type Foxa1 promoter activity but only slightly increased the Foxa1 promoter with mutated AP-1 sites. SUM1315 cells expressing shCtrl or shTwist1 were cultured in 24 well plates and transfected with the indicated Foxa1 promoter-Luc reporters. Luciferase assay was carried out 24 hours later after the transfection. Experiments were repeated 3 times and the representative data with 3 technical replicates are presented as Mean ± SEM. *, P < 0.05 by Student’s t test.

3.11 Twist1-silenced Foxa1 expression is not responsible for Twist1-induced cell morphological change but it mediates the expression of some Twist1-regulated genes

To examine whether Twist1-silenced Foxa1 expression is responsible for Twist1-induced morphology and gene expression changes in breast cancer cells, we stably restored Foxa1 expression in MCF7 Twist1 cell lines (designated as MCF7 Twist1+Foxa1 cell lines) with Twist1-silenced Foxa1 expression (Fig. 3.20A). As expected, MCF7 Ctrl cells
showed epithelial tumor cell morphology identical to their parent MCF7 cells, while MCF7\textsuperscript{Twist1} cells had an elongated spindle shape consistent with their EMT phenotype. MCF7\textsuperscript{Twist1+Foxa1} cells with restored Foxa1 expression showed a cellular morphology very similar to MCF7\textsuperscript{Twist1} cells (Fig. 3.20B), suggesting that the silenced Foxa1 expression is not responsible for Twist1-induced morphological change of MCF7 cells. Furthermore, Twist1 expression effectively repressed E-cadherin, β-catenin, cytokeratin 8 (K8) and ER\textalpha expression in MCF7\textsuperscript{Twist1} cells as compared to MCF7\textsuperscript{Ctrl} cells, while restored Foxa1 expression only partially rescued K8 and ER\textalpha expression in MCF7\textsuperscript{Twist1+Foxa1} cells as compared to MCF7\textsuperscript{Twist1} cells (Fig. 3.20C). Moreover, Twist1 strongly induced vimentin, integrin β1, integrin α5, and MMP9 expression but did not change fibronectin expression in MCF7\textsuperscript{Twist1} cells as compared to MCF7\textsuperscript{Ctrl} cells. Restored Foxa1 expression had no effect on vimentin expression but inhibited Twist1-induced integrin β1, integrin α5 and MMP9 expression and slightly increased fibronectin in MCF7\textsuperscript{Twist1+Foxa1} cells as compared to MCF7\textsuperscript{Twist1} cells (Fig. 3.20 C-F). In addition, Slug (Snail2) expression was increased in MCF7\textsuperscript{Twist1} cells versus MCF7\textsuperscript{Ctrl} cells, but restored Foxa1 expression did not affect Twist1-induced Slug expression in MCF7\textsuperscript{Twist1+Foxa1} cells versus MCF7\textsuperscript{Twist1} cells. The expression levels of Snail1, Zeb1 and Zeb2 mRNAs were very low and remained unchanged in MCF7\textsuperscript{Ctrl}, MCF7\textsuperscript{Twist1} and MCF7\textsuperscript{Twist1+Foxa1} cells. These results suggest that silenced Foxa1 is not responsible for Twist1-induced mesenchymal cell morphology and most EMT marker gene expression, but responsible for permitting the expression of ER\textalpha, K8 and other selective genes related to cell migration and invasion such as integrin β1, integrin α5 and MMP9.
3.12 Silencing Foxa1 expression is required for Twist1 to promote migration, invasion and metastasis in breast cancer cells

Expression of Foxa1 alone showed no effects on MCF7 cell migration and invasion. However, restored Foxa1 expression in MCF7^{Twist1+Foxa1} cells significantly suppressed
Twist1-induced cell migration on the culture plate and invasion through a Matrigel layer (Fig. 3.21 A and B). To examine whether Foxa1 could inhibit Twist1-promoted metastasis in vivo, we injected MCF7<sup>Ctrl</sup>, MCF7<sup>Foxa1</sup>, MCF7<sup>Twist1</sup> and MCF7<sup>Twist1+Foxa1</sup> cells into the tail veins of SCID mice. We examined the lung tissues 4 weeks after injection. We found metastatic tumor foci in 20% (1/5), 0% (0/5), 100% (5/5) and 20% (1/5) of lungs in mice received MCF7<sup>Ctrl</sup>, MCF7<sup>Foxa1</sup>, MCF7<sup>Twist1</sup> and MCF7<sup>Twist1+Foxa1</sup> cells, respectively (Fig. 3.21C). The average number of MCF7<sup>Twist1</sup> cell metastasis foci developed in each lung was significantly increased versus the average numbers of MCF7<sup>Ctrl</sup>, MCF7<sup>Foxa1</sup>, and MCF7<sup>Twist1+Foxa1</sup> cell metastasis foci per lung (Fig. 3.21D). Accordingly, the lung metastasis index reflecting the percentage of tumor area to lung tissue area was more than 25 fold higher in the recipient mice of MCF7<sup>Twist1</sup> cells versus the recipient mice of MCF7<sup>Ctrl</sup> or MCF7<sup>Twist1+Foxa1</sup> cells (Fig. 3.18E). To examine if restored Foxa1 could partially reverse the Twist1-induced basal tumor phenotype in vivo, we injected MCF7<sup>Ctrl</sup>, MCF7<sup>Foxa1</sup>, MCF7<sup>Twist1</sup> and MCF7<sup>Twist1+Foxa1</sup> cells into the mammary gland fat pads of SCID mice to form xenograft tumors and profiled the expression levels of well-established luminal and basal breast cancer marker genes. Luminal markers including FOXA1, PGR, GRP160, BAG1, BLVRA, PDEF, XBP1 and MUC1 were repressed and basal markers including JAG1, EGFR, FOXC1, CK5, CDC20 and ITGB1 were induced in MCF7<sup>Twist1</sup> tumors versus MCF7<sup>Ctrl</sup> and most MCF7<sup>Foxa1</sup> tumors. Restored Foxa1 expression increased the expression of many luminal markers including BLVRA, PDEF and XBP1 and decreased the expression of many basal markers including JAG1, CK5, CDC20 and ITGB1 in MCF7<sup>Twist1+Foxa1</sup> tumors.
versus MCF7<sup>Twist1</sup> tumors (Fig. 3.22A). Again, Twist1 promoted MCF7<sup>Twist1</sup> tumor metastasis, while restored Foxa1 inhibited MCF7<sup>Twist1+Foxa1</sup> tumor metastasis in SCID mice (Fig. 3.22 B and C). Together, these results demonstrate that Twist1-repressed Foxa1 expression may be important for BLBC progression and is required for Twist1-promoted migration, invasion and metastasis of breast cancer cells. These results demonstrate that Twist1-mediated downregulation of Foxa1 expression is required for TWISTS1-promoted breast cancer cell metastasis.

Figure 3.21. Restored Foxa1 expression inhibited Twist1-promoted migration, invasion and metastasis of breast cancer cells. A & B. Cell migration and invasion assays for MCF7 parent, MCF7<sup>Ctrl</sup>, MCF7<sup>Foxa1</sup>, MCF7<sup>Twist1</sup> and MCF7<sup>Twist1+Foxa1</sup> cells. *, P<0.05 by Student’s t test. C. Lung photographs of SCID mice after receiving intravenous injection of MCF7<sup>Ctrl</sup>, MCF7<sup>Foxa1</sup>, MCF7<sup>Twist1</sup> or MCF7<sup>Twist1+Foxa1</sup> cells for 4 weeks. Asterisks indicate visible metastasis nodules. D. Metastasis nodules identified on the H&E-stained serial lung sections of SCID mice (n=5) injected with MCF7<sup>Ctrl</sup>, MCF7<sup>Foxa1</sup>, MCF7<sup>Twist1</sup> or MCF7<sup>Twist1+Foxa1</sup> cells for 4 weeks. **, P<0.01 by Student’s t test. E. Metastasis indexes were presented as the average ratios of tumor areas to lung areas measured on the images taken from H&E-stained serial lung sections. ***, P<0.01 by Student’s t test.
Figure 3.22. Restored Foxa1 expression increased the expression of most Twist1-suppressed luminal marker genes and decreased the expression of most Twist1-promoted basal marker genes in the xenograft tumors in mice. A. MCF7\textsuperscript{Ctrl}, MCF7\textsuperscript{Foa1}, MCF7\textsuperscript{Twist1} or MCF7\textsuperscript{Twist1+Foa1} cells were injected into the second pair mammary gland fat pads of each 8-week-old female SCID mouse (4 × 10\textsuperscript{6}/fat pad). Five mice were used for each group. Injected mice were fed drinking water containing 1µg/ml of 17β-estradiol. In 6 weeks after cell injection, 5, 4, 9 and 7 tumors were observed in mice receiving MCF7\textsuperscript{Ctrl}, MCF7\textsuperscript{Foa1}, MCF7\textsuperscript{Twist1} or MCF7\textsuperscript{Twist1+Foa1} cells, respectively. Mice were euthanized and the xenograft tumors were collected. Total RNA samples were prepared from 4 tumors in each group and qPCR analyses were performed to measure the mRNA levels of the indicated luminal and basal marker genes. Their relative expression levels were obtained by normalized to the level of b-actin mRNA in each sample. Four tumor RNA samples in each group were analyzed in duplicates and data are presented as Mean ± SEM. One-Way ANOVA was performed to analyze the statistical differences among groups. * and **, P < 0.05 and 0.001; NS, not significant. B & C. The lungs were isolated from the above mice injected with MCF7\textsuperscript{Ctrl}, MCF7\textsuperscript{Foa1}, MCF7\textsuperscript{Twist1} or MCF7\textsuperscript{Twist1+Foa1} cells and examined for metastasis by counting the number of metastasis nodules and measuring the ratio of metastatic tumor area to total lung area (Metastasis Index) on the H&E-stained lung sections.
3.13 Breast tumor patients with high Twist1 and low Foxa1 expression exhibit poor distant metastasis-free survival

To assess the clinical relevance of the Twist1-Foxa1 regulatory axis in human breast cancer progression, we analyzed the association between the expression levels of *Twist1* and *Foxa1* and the clinical outcomes in a large breast cancer cohort with distant metastasis-free survival (DMFS) data \(^{121}\). The whole cohort of patients was divided into four groups according to the median *Twist1* and *Foxa1* expression. The *Twist1*\textsuperscript{High}/*Foxa1*\textsuperscript{Low} subgroup exhibited significantly worse DMFS than the *Twist1*\textsuperscript{Low}/*Foxa1*\textsuperscript{High} and the *Twist1*\textsuperscript{High}/*Foxa1*\textsuperscript{High} subgroups. DMFS of the *Twist1*\textsuperscript{Low}/*Foxa1*\textsuperscript{Low} subgroup is not significantly different from the DMFS curves of the other three groups (Fig. 3.23A). Further analyses revealed that the percentages of *Twist1*\textsuperscript{Low}/*Foxa1*\textsuperscript{High} and *Twist1*\textsuperscript{High}/*Foxa1*\textsuperscript{High} subgroup tumors in luminal A and luminal B subtypes are much higher than those in Basal-like and HER2-positive subtypes, while the percentages of *Twist1*\textsuperscript{Low}/*Foxa1*\textsuperscript{Low} and *Twist1*\textsuperscript{High}/*Foxa1*\textsuperscript{Low} tumors in luminal A and luminal B subtypes are much lower than those in the more aggressive basal-like and HER2-positive subtypes (Fig. 3.23B). Interestingly, when the cohort was further
subgrouped into luminal A, luminal B, HER2-positive and basal-like subtypes, only the Twist1\textsuperscript{high}/Foxa1\textsuperscript{Low} patients with luminal A tumors exhibited significantly worse DMFS as compared to Twist1\textsuperscript{Low}/Foxa1\textsuperscript{High} and Twist1\textsuperscript{High}/Foxa1\textsuperscript{High} patients with luminal A tumors (Fig. 3.23 C-F). These results suggest that Twist1-mediated repression of Foxa1 expression plays an important role in breast cancer metastasis and Foxa1 can effectively inhibit metastasis in tumors with high expression of both Twist1 and Foxa1.
Figure 3.23. The association of Twist1 and Foxa1 expression levels with the clinical outcomes of the breast cancer patients. A. The distant metastasis-free survival (DMFS) curves of women with breast tumors expressing high or low levels of Twist1 and Foxa1 as indicated. The cohort (n=1609) was divided into high and low expression subgroups by the median. B. The distribution of patients with the indicated expression patterns of Twist1 and Foxa1 mRNAs in the 4 subtypes of breast cancer. The number of cases in each group is indicated above each bar. C-F. 10-year DMFS curves for luminal A (Panel C), luminal B (Panel D), HER2-positive (Panel E), and basal-like (Panel F) breast cancer patients with different expression levels of Twist1 and Foxa1 mRNAs. Sample number (n) is indicated for each subgroup. **, P<0.01 by Log-rank test.
4. SUMMARY AND DISCUSSION

In this study, we developed the novel TVA/RCIP mouse model system in which deletion of a floxed gene such as Twist1 and expression of an oncogene such as PyMT happen simultaneously in the RCIP virus-infected LECs expressing TVA. This model has several unique advantages. Firstly, TVA is only expressed in the mammary LECs in MMTV-TVA mouse line \(^{105}\) and RCIP virus only infects the TVA-expressing cells, allowing specific delivery of Cre recombinase and PyMT (or any oncogene) to these mammary LECs. Secondly, the intraductally introduced RCIP virus only infects a small population of the mammary LECs expressing TVA and transform these cells into tumor cells. This closely simulates spontaneous tumorigenesis in humans with tumor initiation and growth in a normal surrounding cellular environment. Thirdly, the Cre-coding sequence is located before the short IRES and the PyMT sequences, which makes Cre expression much higher than PyMT expression and guarantees efficient knockout of the floxed genes in the tumor cells. Finally, the single virus construct-mediated co-expressions of Cre and PyMT also save time and resources for crossbreeding of multiple mouse lines. These unique features make this TVA/RCIP model system very useful for characterizing the role for any floxed genes during the process of mammary tumor initiation, progression and metastasis.

Using TVA/RCIP system, we selectively induced mammary tumorigenesis from Twist1 null LECs in mice. We found that Twist1 is not expressed in normal LECs and the RCIP
virus-induced initiation and growth of mammary tumors derived from LECs with or without Twist1 knockout are the same, indicating that Twist1 is not required for mammary tumor formation in LECs. Tumor cell proliferation, apoptosis and macrophage recruitment all showed no significant difference. These results of endogenous Twist1 function further validates the previous studies showing knockdown of Twist1 in breast cancer cell lines did not affect their proliferation in culture and xenograft tumor growth in mice \(^{26,68}\). Interestingly, a recent study reported an essential role of Twist1 in developing skin tumors, suggesting the role of Twist1 in carcinogenesis may depend on cell types \(^{122}\). Another study \(^{123}\) using pancreatic tumor model showed that knock out of Twist1 or Snail1 is dispensable for initiation and progression of primary pancreatic cancer. Our data is consistent with the pancreatic study that Twist1 knockout does not influence breast tumor initiation, proliferation, apoptosis or macrophage recruitment.

Our results showed Twist1 is expressed in a small population of tumor cells at advanced stages at levels similar to the stromal cells, indicating that only certain primary tumor cells gain Twist1 expression during tumor progression. We did not find lung metastasis in mice with palpable mammary primary tumors that have been developed for 6 weeks, but did find extensive lung metastases in mice with palpable mammary primary tumors that have been developed for 10 weeks. These results suggest that Twist1 expression is associated with metastasis in mammary tumors with functional Twist1 gene. More importantly, Twist1 null tumor cells yielded significantly fewer lung metastasis than
tumor cells with functional Twist1, indicating that the functional Twist1 gene promotes tumor cell metastasis. Together, we conclude that Twist1 plays a tumor cell-autonomous function to promote metastasis and validate the results of previous studies showing that knockdown of Twist1 in established breast cancer cell lines inhibits their metastasis in immune-defective host mice 26, 64, 68.

We then examined EMT markers by IF staining and found that Twist1 expressing tumor cells indeed showed EMT marker transition: decreased E-cadherin but increased vimentin, suggesting the association of EMT with lung metastasis. However, because we found metastatic tumor cells in the lung still maintain epithelial phenotype and do not express Twist1, it remains debatable whether the initial Twist1-expressing tumor cells in the mammary glands physically metastasize to the lung. A recent study using a squamous cell carcinoma mouse model with inducible Twist1 expression nicely demonstrated that although Twist1 expression strongly promotes tumor cell invasion and intravasation, turning off Twist1 expression is required for the disseminated tumor cells to establish metastasis in a distant organ 49. According to this model, it is possible that Twist1 expression in primary tumor cells induces partial EMT and drives local invasion and intravasation. Then, Twist expression is turned off in the tumor cells of lung metastases. However, it is also possible that the initial Twist1-expressing tumor cells did not metastasize to the lung, but these cells induce their surrounding Twist1-negative tumor cells to metastasize. Two recently published studies support this model 123, 124.
Since Twist1 is a transcription factor, it should promote breast cancer metastasis through either inhibiting or enhancing its target gene expression. In this study, we found that Twist1 expression negatively correlates with Foxa1 expression in both mouse and human breast tumors. Foxa1 expression is associated with LBC which has a good prognosis. Meanwhile Twist1 is associated with BLBC which has a poor prognosis due to its potential to drive cell invasiveness and metastasis. However, the regulatory and functional relationship between Twist1 and Foxa1 in breast cancer progression remains unknown. In this study, we found Twist1 expression negatively correlates with Foxa1 expression in human breast cancer cells and tumors. Mechanistically, Twist1 directly associates with Foxa1 promoter to recruit NuRD complex and prevent AP-1 binding to Foxa1 promoter, resulting in decreased H3K9 acetylation, reduced RNA polymerase II recruitment and silenced Foxa1 expression. It is conceivable that constitutive AP-1 binding in LBC cells with no Twist1 plays a major role to maintain a high level of Foxa1 expression and thus a luminal phenotype. Twist1 expression in partial or full EMT breast cancer cells may partially or fully inhibit AP-1 recruitment to Foxa1 promoter through NuRD complex-mediated changes in histone codes and thus reduce or silence Foxa1 expression, causing breast cancer progression toward BLBC. Therefore, our findings indicate that Foxa1 is a direct target of Twist1 during breast cancer progression and provides a possible new mechanism for the loss of Foxa1 during breast cancer progression from a luminal ERα-positive subtype to an ERα-negative BLBC subtype.

In agreement with the established function of Twist1 \(^{64}\), ectopic expression of Twist1 in
MCF7 cells induced a mesenchymal morphology and gene expression signature but decreased epithelial signature. Using this cellular model, we addressed how much the Twist1-silenced Foxa1 expression contributes to Twist1-induced mesenchymal morphogenesis and gene expression in breast cancer cells by restoring Foxa1 in Twist1-expressing MCF7 cells. Interestingly, though previous studies have shown Foxa1 represses EMT-linked mesenchymal morphogenesis in pancreatic cancer cells, we found that restored Foxa1 did not change the mesenchymal cell morphology induced by Twist1 in MCF7Twist1+Foxa1 cells. In agreement with this, restored Foxa1 did not rescue the expression level changes of either canonical epithelial genes such as E-cadherin and β-catenin or mesenchymal genes such as vimentin and Slug, which are inhibited or induced by Twist1 expression in MCF7 cells, respectively. We then also estimated the role of Foxa1 silencing in EMT by comparing the genes regulated by Foxa1 in the LBC cells to the previously published EMT signature genes. Among the 1118 Foxa1-regulated genes, we only found ARHGAP8, LMCD1, NMU, PPAP2B, PRKCH and SLPI in the 251 EMT signature gene lists. None of these 6 genes has been shown to regulate the mesenchymal morphogenesis of breast cancer cells. Together, these findings suggest that Twist1-silenced Foxa1 expression is not a major regulatory pathway for the mesenchymal morphogenesis and for the expression of most typical EMT signature genes in breast cancer cells.

Interestingly, restored Foxa1 does partially rescue ERα and K8 expression and strongly downregulate Twist1-induced integrin β1, integrin α5 and MMP9 expression. Since
ERα and K8 are markers of the LBC and integrins β1 and α5 are preferentially expressed in BLBC, it is conceivable that Twist1-silenced Foxa1 expression plays a crucial role in promoting BLBC progression. Foxa1 could either directly or indirectly regulate these genes. It has been reported that Foxa1 upregulates but Twist1 directly represses ERα expression\(^\text{12, 91, 120}\). The results of the present study suggest that both the loss of Foxa1-mediated activation due to Twist1-silenced Foxa1 expression and the Twist1-mediated active repression are required for shutting down ERα expression and developing endocrine resistance during breast cancer progression toward BLBC. These interpretations are consistent with the previous studies showing that Foxa1 expression is associated with good responses to endocrine therapy and the loss of Foxa1 is associated with BLBC progression\(^\text{93, 94, 97-102}\). More importantly, restored Foxa1 robustly inhibited Twist1-promoted migration, invasion and metastasis. These could be partially attributed to the downregulation of Twist1-induced expression of MMP9 and integrins β1 and 5α, since these proteins are known to promote cell invasiveness and metastasis\(^\text{110, 127-131}\). In addition, we demonstrate that Twist1 expressed in MCF7 cells increased most BLBC markers but decreased most LBC markers in these cell-derived xenograft tumors, while restored Foxa1 in Twist1-expressing MCF7 cells decreased many BLBC markers but increased many LBC markers. These results further support the notion that Twist1-repressed Foxa1 expression may play an important role to promote BLBC progression.

In previous studies, the role of Twist1 in promoting metastasis was largely attributed to its capability to induce EMT and enhance cancer stem cell (CSC) features\(^\text{26, 51, 64, 71}\).
Now, the consensus from present and previous studies suggests that Twist1 may serve as a master regulator in breast cancer progression by regulating multiple genes involved in multiple pathways. On one hand, Twist1 may regulate one subgroup of genes, such as downregulating E-cadherin and ERα$^{12, 68, 120}$ and upregulating Bmi1, AKT2, Wnt5a and vimentin $^{70, 85, 132}$ to promote EMT, CSC features, cell migration, invasion and metastasis. On the other hand, Twist1 may regulate another subgroup of genes, such as upregulating PDGFRα to induce invadopodia formation for cell invasion $^{72}$ and downregulating Foxa1 to decrease ERα, CK8 and other LBC markers and increase integrin α5, integrin β1, MMP9 and other BLBC markers for promoting BLBC progression and breast cancer cell migration, invasion and metastasis. Although these multiple Twist1-regulated genes and pathways may not be equally important, they may work cooperatively to drive breast cancer progression and metastasis. A single targeting event such as silencing Foxa1 or upregulating AKT2 can be required but may not be sufficient for Twist1-promoted migration, invasion and metastasis of breast cancer cells.
Analysis of a large cohort dataset showed that tumors with $\text{Twist1}^{\text{High}}\text{Foxa1}^{\text{Low}}$ expression are associated with worse DMFS versus $\text{Twist1}^{\text{Low}}\text{Foxa1}^{\text{High}}$ and $\text{Twist1}^{\text{High}}\text{Foxa1}^{\text{High}}$ tumors, which supports the notion that Twist1-silenced Foxa1 expression can promote breast cancer metastasis. Interestingly, this association is only observed in the mixed subtypes and luminal A subgroup, but not in the luminal B, HER2-positive or basal-like subgroups. We noticed that the tumor numbers with each expression profile are more evenly distributed in the luminal A and luminal B subgroups versus basal-like and HER2-positive subgroups. The latter two subtypes contain too few tumors with $\text{Twist1}^{\text{Low}}\text{Foxa1}^{\text{High}}$ and $\text{Twist1}^{\text{High}}\text{Foxa1}^{\text{High}}$ expressions and thus do not
support a valid statistical analysis to compare Twist1\textsuperscript{Low}Foxa1\textsuperscript{High} or Twist1\textsuperscript{High}Foxa1\textsuperscript{High} tumors with Twist1\textsuperscript{Low}Foxa1\textsuperscript{Low} or Twist1\textsuperscript{High}Foxa1\textsuperscript{Low} tumors. Nevertheless, the facts that \textit{Twist1} is expressed at high levels in more than 40\% of basal-like and HER2-positive tumors and Foxa1 is expressed at low levels in 96\% of basal-like and 77\% of HER2-positive tumors also support the notion that silenced Foxa1 expression is associated with more malignant breast cancer subtypes. For the luminal B subtype group, other factors in addition to Twist1 and Foxa1 may determine DMFS of these patients.

Analysis of tumor tissue microarrays by IHC also revealed an overall negative correlation between Twist1 and Foxa1 proteins. However, we did not found many Twist1-positive tumors from the examined breast tumor microarrays either because of too few BLBCs in the cohort or missed identification of some positive tumors due to the small area of tissue microarrays which are insufficient to cover heterogeneous regions for Twist1 expression in a tumor. IHC revealed that Twist1 is usually expressed only in a small subpopulation of tumor cells or a small region in a tumor. Most of these Twist1-positive tumors also contain many Foxa1-positive luminal tumor cells. These double positive tumors may contain Twist1-positive/Foxa1-positive, Twist1-positive/Foxa1-negative and/or Twist1-negative/Foxa1-positive cells. These observations support the notion that Twist1 is expressed in a subset of breast tumor cells and that Twist1 repressed Foxa1 expression and drive EMT to promote BLBC progression from this subset of tumor cells.
In summary, we found that endogenous Twist1 mainly promotes breast cancer invasiveness and metastasis with no obvious function in affecting primary tumor development. Twist1 is expressed in a small population of tumor cells and its expression associates with EMT phenotype, invasiveness and metastasis. We further found that Twist1 directly represses the transcriptional activity of the Foxa1 promoter in breast cancer cells. Twist1-silenced Foxa1 expression is largely responsible for Twist1-mediated migration, invasion and metastasis but less responsible for Twist1-induced EMT morphology. On the other hand, silencing Foxa1 by Twist1 in breast cancer strongly promotes the transition from the luminal to basal gene signatures. These results not only uncover the mechanism of Foxa1 silencing in BLBC but also provide a new mechanism of Twist1’s role in BLBC progression. Therefore, targeting Twist1 in BLBC treatment may be an effective approach to inhibit invasion and metastasis.
REFERENCES


