SHORT COMMUNICATION

Isoprenylcysteine carboxylmethyltransferase is critical for malignant transformation and tumor maintenance by all RAS isoforms

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Despite extensive effort, there has been limited progress in the development of direct RAS inhibitors. Targeting isoprenylcysteine carboxylmethyltransferase (ICMT), a unique enzyme of RAS post-translational modification, represents a promising strategy to inhibit RAS function. However, there lacks direct genetic evidence on the role of ICMT in RAS-driven human cancer initiation and maintenance. Using CRISPR/Cas9 genome editing, we have created Icmt loss-of-function isogenic cell lines for both RAS-transformed human mammary epithelial cells (HME1) and human cancer cell lines MiaPaca-2 and MDA-MB-231 containing naturally occurring mutant KRAS. In both in vitro and in vivo tumorigenesis studies, Icmt loss-of-function abolishes the tumor initiation ability of all major isoforms of mutant RAS in HME1 cells, and the tumor maintenance capacity of MiaPaca-2 and MDA-MB-231 cells, establishing the critical role of ICMT in RAS-driven cancers.

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INTRODUCTION

Constitutively active mutations of RAS are the most common driving mutations in human cancers,1,2 with up to 90% of invasive pancreatic ductal adenocarcinomas harboring activating KRAS mutations. Due to the high prevalence and established causal role of RAS in tumorigenesis and tumor progression, there have been remarkable efforts on the therapeutic targeting of RAS,3 albeit with only limited progress.4,5 Effective therapy for mutant RAS-driven tumorigenesis therefore remains an unresolved challenge.

Inhibiting RAS-driven processes by targeting RAS trafficking and membrane localization has been an important strategy to circumvent the difficulties in targeting RAS directly.1,4–7 Most efforts in this regard have focused on targeting the post-translational processing of RAS proteins. RAS proteins carry the so-called CaaX motif at their C-terminus that subjects them to sequential prenylation, endoproteolytic cleavage and carboxymethylation.1,6 Isoprenylcysteine carboxylmethyltransferase (ICMT), the enzyme catalyzing the carboxymethylation step,8 has garnered attention in recent years because it is the only enzyme known to catalyze this important modification, in contrast to the two protein prenyltransferases catalyzing the first step of the processing.6,9,10 therefore lowering the risk of target failure. Recent studies have shown that small-molecule inhibitors of ICMT have good efficacy against several human cancer cell lines11,12 possibly through an impact on energy metabolism and autophagy,13 and display synergistic activity with other antitumor agents.14,15

Mouse genetic studies have supported the importance of ICMT in RAS localization and function16; however, these models have provided some conflicting results on the role of ICMT on RAS-driven tumorigenesis in mouse.17,18 There is also a lack of direct genetic evidence in human cancer cells on the role of ICMT in RAS-driven transformation and maintenance. Another important unanswered question in this field is whether ICMT inhibition impacts tumorigenic processes driven by different RAS isoforms, since most studies have been done on KRAS up to this point.

In this study, we used CRISPR/Cas9 genomic editing to create Icmt loss-of-function isogenic cell lines to investigate the role of ICMT in RAS-driven tumor initiation and maintenance in two systems. In one approach, we employed a stepwise-transformed human mammary epithelial cell (HME1) model to investigate the impact of ICMT function on tumorigenesis driven by different RAS isoforms. In the second approach, MiaPaca-2 and MDA-MB-231, which are KRAS mutant patient-derived aggressive human pancreatic and breast cancer cell lines, respectively, were used to generate Icmt loss-of-function isogenic cells to study the impact of ICMT on tumor maintenance, in both in vitro and in vivo systems.

RESULTS AND DISCUSSION

HME1 cells can be efficiently transformed by G12V-mutant, but not wild-type, RAS isoforms.

To construct a model system to study the role of ICMT in RAS-driven epithelial tumorigenesis, we obtained immortalized HME1 cells stably expressing hTERT and generated, stepwise, isogenic cell lines expressing SV40 small T antigen and a shRNA against p53 using retroviral vectors; the resultant line is termed

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Figure 1. HME1-shp53 cells can be transformed by all four RAS-G12V mutant isoforms, but not by their wild-type counterparts. (a) Illustration of the scheme for creation of the HME1-shp53 cell line by sequential addition of SV40 small T antigen, shRNA against p53, and ER-tagged wild-type or G12V-mutant RAS isoforms into HME1-hTERT cells. The parental HME1-hTERT (ATCC CRL-4010™) cells were obtained from American Type Cell Culture (ATCC) and mycoplasma-free. Stable expression of genes was achieved by transduction with retroviral expression vectors pBABE-hygro (for SV40 Small T Antigen), pRetroSuper (for shRNA against p53) or pBABE-puro (for all forms of RAS) with viruses generated via calcium phosphate transfection, followed by stable selection using 20 μg/ml hygromycin for pBABE-hygro (Invitrogen, Carlsbad, CA, USA), 5 μg/ml blasticidin for pRetroSuper (Sigma Aldrich, St Louis, MO, USA), and 5 μg/ml puromycin (Sigma Aldrich) for pBABE-puro. Ras isoforms were cloned using primers listed in Supplementary Table S1. G12V mutations were introduced using QuikChange Site-Directed Mutagenesis Kit (Agilent Technologies) and the targeting oligonucleotides are listed in Supplementary Table S1. (b) HME1-shp53 cells expressing wild-type or ER-tagged G12V mutant of KRAS4A, KRAS4B, HRAS and NRAS as shown in (a) were cultured in soft agar in the presence or absence of 500 nM tamoxifen (Sigma Aldrich) for 14 days, and subsequently stained using methylthiazolyldiphenyl-tetrazolium bromide (MTT; Sigma Aldrich). This three-dimensional anchorage-independent soft agar culture was performed according to a previously published protocol supplemented with 0.5 μg/ml hydrocortisone (Sigma Aldrich), and 5 μg/ml insulin (I-DNA Biotech, Singapore). Using an Olympus SZX16 stereo microscope with 3.2 × zoom and 0.5 × objective, pictures of 4–5 random microscopy fields were taken per well after MTT staining; one representative field for each condition from one representative experiment out of three performed is presented. (c) Immunoblot study of cell lysates from HME1-shp53 control cells and those expressing RAS proteins. The cell lysates were prepared from the same cells used for the soft agar study in (b) and HME1-shp53 was used as the negative control. Immunoblots were done on cell lysates per standard protocol established in the laboratory. The antibodies for pan-RAS (Ras10, MA1-012) and GAPDH (14C10, #2118) were from Pierce Scientific (Waltham, MA, USA) and Cell Signaling Technology (Danver, MA, USA), respectively. Results presented in (b) and (c) are from a single experiment that is representative of three similarly performed studies.
HME1-shp53 (Figure 1a). Subsequently, we investigated the ability of the major isoforms of wild-type and mutant (G12V) RAS proteins to transform these epithelial cells by introducing retroviruses that express estrogen receptor (ER)-tagged RAS isoforms to create eight HME1-shp53-ER-RAS stable cell lines (Figure 1a).

These cells were then cultured in soft agar in the presence or absence of tamoxifen to investigate their ability to grow colonies under anchorage-independent conditions. In the absence of tamoxifen treatment, ER-RAS proteins were minimally detected in all four lines (Figure 1c), and soft agar colony

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**Figure 2.** For Caption see next page.
formation was not evident (Figure 1b). Tamoxifen treatment induced ER-RAS protein expression markedly for all wild-type and mutant RAS isoforms, with some variation in expression levels between different isoforms (Figure 1c). Importantly, robust colony growth was observed for cells expressing all RAS-G12V isoforms, while there was no colony formation from cells expressing wild-type isoforms (Figure 1b), despite comparable RAS expression levels induced by tamoxifen treatment between the wild type and mutant for each isoform (Figure 1c). The establishment of the HME1 system that can achieve efficient transformation by all G12V-mutant RAS isoforms allowed us to further investigate the role of ICMT in tumorigenesis elicited by mutant RAS.

Loss of ICMT function abolishes the transforming ability of all four RAS-G12V isoforms

To evaluate the role of ICMT in RAS-G12V-driven HMEC transformation, the CRISPR/Cas9 method was used to generate loss-of-function of ICMT from HME1-shp53 cells (Figure 2a). The loss of both copies of the functional Icmt coding sequence was confirmed by Sanger sequencing, which showed frameshift mutations that would translate into mutant truncated polypeptides (Figure 2b, Supplementary Table S2). Several confirmed clones expressing each truncated polypeptide (1cr and 4cr) were combined in equal number to generate the two ICMT loss-of-function cell lines (HME1-shp53 ICMT-null 1cr and HME1-shp53 ICMT-null 4cr; Figures 2a and b) that contain only the very N-terminal portion of the ICMT polypeptide, as shown in Figure 2b, which would retain no ICMT function.21 Similarly, the HME1-shp53 wild-type Icmt cell line was a mix of three clones arising from HME1-shp53 cells transfected with empty vector. Subsequently, the retroviral vector pBabe-puro was used to express the four ER-tagged RAS-G12V proteins in the three cell lines generated (Figure 2a). As noted above, this inducible system enabled us to observe RAS-G12V-driven transformation without the concern of potential signaling adaptation associated with constitutive expression of mutant-RAS. Another advantage of this system is that the RAS expression level can be manipulated by titrating the concentration of tamoxifen; in this way, the exogenous expression can be controlled to be near physiological levels.

We found that inducing KRAS, NRAS and HRAS with 500, 100 and 25 nM of tamoxifen, respectively, allowed us to achieve expression of the RAS isoforms at levels comparable to those in cancer cell lines known to harbor activating mutations of RAS.

Figure 2. Eliminating ICMT function abolishes the transforming ability of G12V-mutant RAS. (a) Schematic illustration showing the creation of HME1-shp53 ICMT-wild type, ICMT-null 1cr and ICMT-null 4cr cells. Wild-type ICMT expression was eliminated from HME1-shp53 by creating frameshift mutations in the genomic Icmt loci using the CRISPR/Cas9 method via expression vector pSpCas9 (BB)-2A-GFP (Addgene #48130).21 Control empty vector pSpCas9 (BB)-2A-GFP (EV), as well as the same vector carrying guide RNA sequences (termed 1cr and 4cr, Supplementary Table S2) were introduced into cells by electroporation using Gene Pulser Xcell™ (Bio-Rad, Hercules, CA, USA). Single cells expressing GFP were individually sorted using BD FACSAria™ III (BD Biosciences, Franklin Lakes, NJ, USA) into 96-well plates for clonal expansion and subsequently screened for genomic frameshift mutations by fluorescent PCR-capillary gel electrophoresis using a protocol by Ramlee et al.21 Standard PCR and Sanger sequencing were performed to identify the presence of frameshift mutations in exon 1. (b) Schematic illustration of the predicted polypeptide translated from each allele of HME-1 shp53 cells expressing wild-type ICMT,21 as well as of truncated polypeptides translated from mutant alleles in individual HME-1 shp53 ICMT-null clones. Green, cytoplasmic motifs; pink, endoplasmic reticulum luminal motifs; yellow, transmembrane domains; white, frameshifted nonsense polypeptide sequences resulting from CRISPR/Cas9 action; asterisk (*), end of polypeptide chain. The sequences of the mutated alleles are shown in Supplementary Table S2. (c) Immunoblot study of lysates of HME1-shp53-ICMT wild type, ICMT-null 1cr, and ICMT-null 4cr cells expressing ER-tagged G12V-mutant KRAS4A, KRAS4B, HRAS or NRAS. These cells were cultured in the presence or absence of tamoxifen for 48 h, then harvested for lysate preparation for immunoblot analysis. Lysates of cancer cell lines (MiaPaCa-2, MDA-MB-231, NCI-H1299 and T24; HRAS; Figure 2c). The discrepancies in expression level among different ER-RAS isoforms are consistent with data suggesting that differences in rare codon utilization in the coding sequences significantly affect the expression levels of RAS isoforms.22 We then evaluated the transformation capacity of all four RAS-G12V isoforms in the presence and absence of functional ICMT using the soft agar colony formation assay. As expected, HME1-shp53 cells in which no ER-RAS coding sequence was introduced (empty retroviral vector) did not form colonies with tamoxifen concentration at the highest level of 500 nM regardless of ICMT status, while robust colony formation was observed with ICMT-wild-type cells expressing ER-tagged G12V mutants of KRAS4A, KRAS4B, NRAS and HRAS in the presence of tamoxifen (Figure 2d). Importantly, the loss of ICMT function in the cell lines harboring all four RAS-G12V isoforms resulted in abrogation of tamoxifen-induced colony formation ability (Figure 2d) without affecting the level of ER-RAS proteins (Figure 2c). These data indicate that the transforming capacity of all four isoforms of RAS is sensitive to the impairment of their C-terminal methylation.

Loss of ICMT function abrogates the formation of xenograft tumors by cells harboring mutant RAS

To investigate the relevance of ICMT in RAS-driven tumorigenesis in vivo, we created a set of ICMT-wild-type and ICMT loss-of-function HME1 cells constitutively expressing NRAS-G12V and KRAS4B-G12V without the ER tag to circumvent the problems associated with dosing mice long term with tamoxifen. Two ICMT loss-of-function cell lines (HME1-shp53 ICMT-null 1cr and HME1-shp53 ICMT-null 4cr) and the control ICMT-wild-type cell line (HME1-shp53 ICMT-wild type; see above) were infected with retroviruses expressing constitutively active NRAS-G12V and KRAS4B-G12V isoforms. Exogenous NRAS and KRAS expression was confirmed in the respective cell lines through quantitative real-time PCR (Supplementary Figure S1). As expected, in soft agar assays minimal colonies were formed from cells infected only with the empty vector, regardless of their ICMT status. In contrast, HME1-shp53 cells were successfully transformed by constitutively active NRAS-G12V and KRAS4B-G12V in cells with functional ICMT, as demonstrated by the robust anchorage-independent colony growth in soft agar (Figure 3a). Moreover, loss of ICMT function in both the ICMT-null 1cr and ICMT-null 4cr cell lines dramatically reduced the colony growth elicited by expression of constitutively active mutant NRAS and KRAS in the...
Importantly, re-introduction of ICMT by retroviral expression of GFP-tagged ICMT reinstated the colony-forming capacity of these cells (Figure 3a). Immunoblot analysis of prelamin A showed the expected accumulation in cells lacking ICMT function, and the prelamin A levels reverted to baseline when ICMT was re-introduced in these cells (Figure 3b). To assess tumorigenesis in vivo, $5 \times 10^6$ of HME1-shp53-NRAS G12V-ICMT-wild type, -ICMT-null 1cr or -ICMT-null 4cr cells were implanted subcutaneously in the bilateral flanks in SCID mice. The ICMT-wild-type cells and ICMT-null cells were placed in opposite flanks of the mouse. HME1-shp53-ICMT-null 1cr and ICMT-null 4cr cells expressing constitutively active NRAS-G12V failed to form tumors up to 80 days after implantation, whereas the ICMT-wild-type NRAS-G12V-expressing cells initiated robust tumors following implantation (Figures 3c–f). The abilities of HME1-shp53-CA-NRAS4A

![Immunoblot analysis of prelamin A, pan-RAS, and GAPDH in cells with different ICMT statuses](image)

![Graphs showing tumor size over time for different conditions](image)
G12V-ICMT-wild-type, and associated ICMT-null 1cr and ICMT-null 4cr cells, to form xenograft tumors were also studied. Only ICMT-wild-type KRAS4A-G12V cells, but not the corresponding ICMT-null cells, formed tumors, albeit with a longer latency period than those expressing NRAS-G12V (Figures 3g–j).

ICMT function is critical for the tumorigenesis capacity of naturally occurring mutant KRAS human cancer cells

While the HME1 stepwise-transformation model provided a clean background to study the role of ICMT in epithelial cell transformation driven by mutant RAS, argon treatments can be made in regard to whether the same conclusions can be extended to human cancers driven by mutant RAS, which would undoubtedly have more genetic alterations and heterogeneity. To address these questions, ICMT loss-of-function isogenic lines were created, using a similar CRISPR/Cas9 method, from the pancreatic cancer cell line MiaPaCa-2 and breast cancer cell line MDA-MB-231; these cancer cell lines carry KRAS-G12C and KRAS-G13D activating mutations, respectively. Worth noting, the generation of ICMT loss-of-function cell lines was more complicated in cancer cells, since most contain more than two copies of the gene loci; for example, MiaPaCa-2 and MDA-MB-231 cells both contain three copies of ICMT loci. Three exon 1-targeting guide RNA sequences, designated 1cr, 3cr, and 4cr, were selected for prenylating clones of ICMT-null cells, confirmed by Sanger sequencing, for each of the cancer cell lines (Figures 4a and f, Supplementary Table S2). The ICMT-null status of the engineered cancer cell lines was further confirmed by prelamin A accumulation with respect to ICMT-wild-type cells (Supplementary Figure S2). These isogenic clones were introduced into SCID mice in similar manners as the HME1 cells above, that is, ICMT-wild-type cells and null cells were implanted on the contra-lateral flanks. MiaPaCa-2 cells selected for control vector (ICMT-wild-type) formed tumors of approximately 2000 mm³ within 40 days of implantation with a latency of 10 days, while minimal to no growth was detected at the time of euthanization for the cells in which Icmt had been inactivated (Figures 4b–e). Similar results were observed for MDA-MB-231 isogenic cell xenografts (Figures 4g–j). Together, these results demonstrate that ICMT function is critical for the initiation and maintenance of mutant KRAS-driven human cancers.

The development of inhibitors against the enzymes involved in post-translational modification of RAS is considered a promising approach for the treatment of RAS-driven cancers. ICMT, the final of the three enzymes in the prenylation cascade for RAS, has been a focus of therapy development, as it is a unique enzyme not displaying functional redundancy with another enzyme, unlike the two CAAX protein prenyltransferases.6 There is mounting evidence indicating that ICMT is a viable therapeutic target in cancers harboring activating RAS mutations.11,12,14,16,17,25 However, there are significant gaps in the understanding of the role of ICMT in RAS-driven tumorigenesis and tumor maintenance that need to be addressed for further development of ICMT inhibitors for cancer therapy. One of the biggest challenges has been the lack of direct genetic validation of the functional roles of ICMT in RAS-driven tumorigenesis in human cancers, and the identification of specific signaling perturbation(s) conferred by the manipulation of ICMT.

Previous studies on ICMT involvement in human cancer cells have mostly involved the use of pharmacological inhibitors of ICMT,11–14,25 which would evoke the consideration of potential off-target effects that could confound conclusions of specificity and efficacy. Up to now, models of Icmt genetic manipulation have all been developmental mouse models in mice, with Icmt knockout and mutant RAS activation occurring concurrently in a specific tissue early in development.17,18 In these models, changes in the signaling landscape resulting from the loss of ICMT early in development could potentially confound the role of ICMT in RAS-driven tumors that develop later in life. Further, these types of loss-of-function mouse models do not mimic the therapeutic targeting of oncoproteins post malignant transformation, thus restricting their utility in understanding how ICMT manipulation would affect tumor growth and maintenance in human cancers. To build model systems of human cancers that can be used to study the role of ICMT in RAS tumorigenesis, we capitalized on the recent advances of CRISPR/Cas9 genomic editing and developed a model system of pre-malignant human mammary epithelial cells (HME1-shp53) with or without functional ICMT. The studies with these cells subsequently transformed by mutant RAS provided direct evidence that the function of ICMT is critical in the transformation of human epithelial cells by all major isoforms of mutant RAS.

While the HME1 system is an excellent way to assess oncogenesis in a genetically defined epithelial cell type, it does not fully recapitulate the situation in naturally occurring human cancers, which typically contain a much larger number of genetic alterations. Hence, we felt it important to also introduce ICMT loss-of-function alleles into cell lines derived from naturally occurring human cancers, which would allow us to evaluate the role of ICMT in tumor formation and maintenance in more complex cancer models.
Transformation by all RAS isoforms requires ICMT

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cells. Using the isogenic lines created by the CRISPR/Cas9 method from two aggressive human cancer cell lines harboring KRAS mutations—the pancreatic cancer cell line Mia-PaCa2 and breast cancer cell line MDA-MD-231—we confirmed the critical role of ICMT in the development and maintenance of the malignant phenotype. The impact of ICMT inhibition in these cancer cell lines demonstrated that, even on the complex genetic background, RAS-driven cancers can be targeted by ICMT inhibition. There are
remaining questions, of course, in particular as to whether the antitumor impact of ICMT inhibition is solely due to its effect on RAS, and whether RAS-driven cancers with additional oncogenic perturbations can be effectively suppressed by ICMT inhibition. It is imperative, therefore, to better understand the signaling events regulated by ICMT that are essential for malignancies; such information would facilitate identification of the target populations of cancers for effective therapeutic ICMT inhibition. The human ICMT loss-of-function cell models described here should be useful tools in this quest.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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