The Initiator Methionine tRNA Drives Secretion of Type II Collagen from Stromal Fibroblasts to Promote Tumor Growth and Angiogenesis

Highlights
- A mechanistic link between the tRNA repertoire and cancer progression
- Selective control of the secretome by levels of the initiator methionine tRNA
- A link between the tRNAome and ECM secretion from stromal fibroblasts
- A new transgenic model recapitulating a key aspect of the cancer tRNAome

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In Brief
Clarke et al. show that levels of the initiator tRNA\textsubscript{Met} are increased in carcinoma-associated fibroblasts. Using a transgenic approach to recapitulate this in vivo and quantitative mass spectrometry to characterize the stromal secretome, they show that elevated tRNA\textsubscript{Met} drives production of a type II collagen-rich ECM to drive tumor progression.
The Initiator Methionine tRNA Drives Secretion of Type II Collagen from Stromal Fibroblasts to Promote Tumor Growth and Angiogenesis

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SUMMARY

Expression of the initiator methionine tRNA (tRNA<sub>Met</sub>) is deregulated in cancer. Despite this fact, it is not currently known how tRNA<sub>Met</sub> expression levels influence tumor progression. We have found that tRNA<sub>Met</sub> expression is increased in carcinoma-associated fibroblasts, implicating deregulated expression of tRNA<sub>Met</sub> in the tumor stroma as a possible contributor to tumor progression. To investigate how elevated stromal tRNA<sub>Met</sub> contributes to tumor progression, we generated a mouse expressing additional copies of the tRNA<sub>Met</sub> gene (2+tRNA<sub>Met</sub> mouse). Growth and vascularization of subcutaneous tumor allografts was enhanced in 2+tRNA<sub>Met</sub> mice compared with wild-type littermate controls. Extracellular matrix (ECM) deposited by fibroblasts from 2+tRNA<sub>Met</sub> mice supported enhanced endothelial cell and fibroblast migration. SILAC mass spectrometry indicated that elevated expression of tRNA<sub>Met</sub> significantly increased synthesis and secretion of certain types of collagen, in particular type II collagen. Suppression of type II collagen opposed the ability of tRNA<sub>Met</sub> overexpressing fibroblasts to deposit pro-migratory ECM. We used the prolyl hydroxylase inhibitor ethyl-3,4-dihydroxybenzoate (DHB) to determine whether collagen synthesis contributes to the tRNA<sub>Met</sub>-driven pro-tumorigenic stroma in vivo. DHB had no effect on the growth of syngeneic allografts in wild-type mice but opposed the ability of 2+tRNA<sub>Met</sub> mice to support increased angiogenesis and tumor growth. Finally, collagen II expression predicts poor prognosis in high-grade serous ovarian carcinoma. Taken together, these data indicate that increased tRNA<sub>Met</sub> levels contribute to tumor progression by enhancing the ability of stromal fibroblasts to synthesize and secrete a type II collagen-rich ECM that supports endothelial cell migration and angiogenesis.

INTRODUCTION

The use of transcriptomics and proteomics to understand cellular function has been pivotal in advancing our knowledge of the molecular basis of disease processes, and the combined use of these approaches is beginning to highlight discord between the control of gene expression at the levels of transcription and translation. Indeed, implementation of numerous post-transcriptional processes means that expression of particular genes at the mRNA level does not necessarily correlate with cellular concentrations of functional products of the same transcripts. tRNAs are one of the most-abundant RNAs in eukaryotic cells, yet are frequently disregarded in most genome-wide analyses because they are assumed to be background genes with merely housekeeping functions. However, it is now becoming clear that cellular levels of tRNAs are key to the control of gene expression in a number of different contexts. Indeed, tissue-specific differences in tRNA expression occur across a variety of human tissues, with relative levels of tRNAs correlating to the codon usage of highly expressed genes in those tissues [1]. Indeed, the availability of certain tRNAs may be controlled in the cell in order to regulate the efficiency of translation of certain gene transcripts. This may, in turn, allow the cell to regulate the activity of certain pathways [2]. For instance, proliferating cells have increased levels of tRNAs that are specific for codons overrepresented in pathways required for proliferation. A number of publications have demonstrated that tRNA levels may be increased in cancer [3–5], and although it was previously unknown whether this was a cause or consequence of carcinogenesis, recent work from
Gingold et al. [2] highlighted the possibility that tRNA expression could indeed play a role in carcinogenesis itself.

There has also been some focus on the consequences of altered expression of specific tRNAs, including tRNA^{Met}, the initiator methionine tRNA responsible for recognizing the start codon and initiating translation. Whereas one group indicated that overexpression of tRNA^{Met} may reprogram global tRNA expression levels and increase proliferation of human epithelial cells [6], another showed that increased levels of tRNA^{Met} influence the growth of Drosophila larvae largely owing to increased secretion of insulin-related peptides from fat bodies [7]. This in vivo observation that tRNA^{Met} levels can drive protein secretion is particularly interesting. It indicates that tRNA^{Met}-sensitive translational control may contribute to cancer progression not only by altering gene expression of the cancer cells themselves but also by altering the secretome of other cells that support carcinogenesis [8–10].

In the current study, we find that levels of tRNA^{Met} are increased in cancer-associated fibroblasts (CAFs) by comparison to normal fibroblasts. To further understand the consequences of altered stromal expression of tRNA^{Met}, we generated a transgenic mouse that carries two additional copies of the tRNA^{Met} gene (2+tRNA^{Met} mouse). We found that this leads to enhanced tumor growth and increased tumor vascularization. In terms of mechanism, we show that increased tumor growth in 2+tRNA^{Met} mice is due to elevated secretion of type II collagen by fibroblasts which, in turn, supports migration of endothelial cells, leading to increased tumor vascularity. Our data therefore provide the first evidence that tRNA^{Met} levels selectively drive secretion of specific components of the extracellular matrix (ECM) to support tumor progression.

**RESULTS**

**Increased Expression of tRNA^{Met} in the Host Animal Promotes Tumor Growth and Angiogenesis**

To determine whether tRNA expression is altered in the tumor stroma, we measured tRNA levels in immortalized human mammary carcinoma-associated fibroblasts (iCAFs) and compared these with immortalized matched normal fibroblast (iNF) controls. iCAFs were generated by co-injecting INFs with Ras-transformed MCF7 cells subcutaneously into mice [11]. These iCAFs are established to have an enhanced capacity to promote tumor growth and angiogenesis when co-injected as xenografts with a variety of tumor cell types and are established to maintain their myofibroblast-like phenotype when grown in culture [11]. We characterized these cells for the expression of non-coding RNA transcribed by RNA polymerase III. Importantly, increased expression of tRNA^{Met} and the tRNA for isoleucine (tRNA^{Ile}) was seen in iCAFs by comparison with INFs, whereas no change was detected in the expression of the 5S tRNA, another non-coding RNA transcribed by RNA polymerase III (Figure S1).

To investigate the consequences of increased stromal tRNA^{Met}, we generated a transgenic mouse that carries two extra copies of the tRNA^{Met} gene inserted into the genome at the HPRT locus (2+tRNA^{Met} mouse; Figure S2A). 2+tRNA^{Met} animals have elevated levels (1.3- to 1.5-fold) of tRNA^{Met} in a number of tissues, including fibroblasts from lung and from whole-mouse embryos (MEFs) (Figure S2B). 2+tRNA^{Met} animals were the same size and weight as their littermate controls (not shown), and MEFs isolated from 2+tRNA^{Met} embryos did not have altered rates of cellular protein synthesis (measured by 35S-methionine incorporation) or cell proliferation (Figures S2C and S2D). We introduced syngeneic tumor cells into 2+tRNA^{Met} mice as subcutaneous allografts and recorded their growth. Initially, we used transformed melanoblasts derived from Tyr::NrasG61K/::INK4a−/− mice. These transformed melanoblasts are null for INK4a, express mutant NRas under a melanoblast-specific promoter, and form melanomas in syngeneic host mice [12]. Mice were culled once tumors reached a specified size (15 × 15 mm), which we defined as the clinical endpoint in order to perform Kaplan-Meier survival analysis. Importantly, tumors reached clinical endpoint significantly faster in 2+tRNA^{Met} mice than they did in wild-type littermate controls (Figure 1A).

To confirm this observation of enhanced tumor growth in 2+tRNA^{Met} mice, we also measured the growth of Lewis lung carcinoma (LLC) and B16 F0 melanoma allografts. Both allograft models formed significantly larger and more-vascularized tumors in 2+tRNA^{Met} mice than they did when grown in wild-type littermate control animals (Figures 1B and 1C), indicating that increased tRNA^{Met} expression in the host animal can promote increased tumor growth and tumor angiogenesis.

**Increased Stromal tRNA^{Met} Expression Promotes the Deposition of a Pro-migratory ECM via Release of Secreted Factors**

A key component of the tumor microenvironment that dictates cancer cell behavior is the ECM deposited by stromal fibroblasts. To determine whether tRNA^{Met} levels influence the characteristics of fibroblast-derived ECM, we isolated fibroblasts from the lungs and embryos of 2+tRNA^{Met} mice (Figure S2B) and used these to prepare cell-derived matrices using established methods [13, 14]. We then investigated the capacity of this cell-free ECM to support the migration of other cells. ECM deposited by primary cultured fibroblasts from 2+tRNA^{Met} mice had significantly enhanced ability to support migration of both endothelial cells (Figure 2A) and fibroblasts (Figure 2B) by comparison with ECM deposited by cells from their wild-type littermate controls. As a tool to determine how increased tRNA^{Met} expression influences ECM deposition, we generated independent pools of immortalized mouse embryonic fibroblasts (iMEF) transfected with either tRNA^{Met} (iMEF-tRNA^{Met}) or an empty vector control (iMEF-vector) (Figure S3A). Consistent with our observations in primary cultured MEFs, increased tRNA^{Met} expression did not lead to increased cell proliferation or cellular protein synthesis in immortalized MEFs (Figures S3B and S3C). However, ECM deposited by iMEF-tRNA^{Met} cells had a similarly enhanced ability to support increased cell migration as did the ECM secreted by primary cultured fibroblasts from 2+tRNA^{Met} mice (Figure 3A).

To determine whether the relationship between tRNA^{Met} expression and the deposition of pro-migratory ECM was observable in fibroblasts from human tumors, we obtained primary CAFs from two human breast tumors and compared these with fibroblasts isolated from the surrounding normal tissue (>5 cm from the tumor) from the same patients. The CAFs from patient 1 displayed significant upregulation of tRNA^{Met} by comparison with fibroblasts from surrounding normal tissue,
shows final tumor volume ± SEM. The graph on the left shows tumor vessel vessels counted across the entire tumor section. The graph on the right fixed and stained to visualize endomucin for blood vessel identification and then tumors were harvested 22 days from this time point. Tumors were injected subcutaneously into either wild-type (WT) or 2+tRNAi

\[ \text{log rank (Mantel-Cox) test; } *p < 0.05. \]

The graph on the right shows final tumor volume ± SEM. Tumors were harvested 22 days from this time point. Tumors were injected subcutaneously into either wild-type (WT) or 2+tRNAi

\[ \text{log rank (Mantel-Cox) test; } *p < 0.05. \]

Figure 1. Increased Expression of tRNA_{Met} in the Host Animal Promotes Angiogenesis and Growth of Allografts

(A) \(1 \times 10^6\) transformed melanocytes (Tyr::NrasQ61K/\(^{-}\);tNK4a/\(^{-}\)) were injected subcutaneously into either wild-type (WT) or 2+tRNA_{Met} transgenic male mice and tumor size monitored. Mice were culled once tumors reached endpoint (15 x 15 mm in size). Kaplan-Meier survival curves were plotted based on time taken to reach endpoint. n = 14 (WT); n = 14 (2+tRNA_{Met} mice); log rank (Mantel-Cox) test; *p < 0.05.

(B) \(1 \times 10^6\) LLC cells were injected subcutaneously into either wild-type (WT) or 2+tRNA_{Met} transgenic male mice, and tumors were harvested at 21 days. The graph on the right shows final tumor volume ± SEM. Tumors were fixed and stained to visualize endomucin for blood vessel identification and vessels counted across the entire tumor section. Graph on the left shows tumor vessel density ± SEM; n = 10 (WT); n = 10 (2+tRNA_{Met}); unpaired t test; *p < 0.05.

(C) \(0.5 \times 10^6\) B16 F0 mouse melanoma cells were injected subcutaneously into either wild-type (WT) or 2+tRNA_{Met} transgenic mice. The time at which palpable tumors were formed was recorded (minimum size of 3 x 3 mm) and then tumors were harvested 22 days from this time point. Tumors were fixed and stained to visualize endomucin for blood vessel identification and vessels counted across the entire tumor section. The graph on the right shows final tumor volume ± SEM. The graph on the left shows tumor vessel density ± SEM; n = 20 (WT); n = 20 (2+tRNA_{Met}); unpaired t test; *p < 0.05; **p < 0.005.

but this was not observed in the second patient (patient 2) (Figure S4A). Interestingly, the ECM produced by CAFs from patient 1 supported significantly increased cell migration, whereas the CAF-derived ECM from patient 2 did not (Figure S4B). These data indicate that, although upregulation of tRNA_{Met} in CAFs varies between patients, the relationship between tRNA_{Met} levels and the ability to generate pro-migratory ECM may be observed in primary cultured CAFs from human tumors.

Many ECM components are initially secreted by fibroblasts as soluble proteins that are subsequently incorporated into the insoluble ECM. To determine which of these stages of ECM deposition are influenced by tRNA_{Met} levels, we collected conditioned medium from iMEF-tRNA_{Met} and iMEF-vector cells and incubated these with normal iMEFs, which were, in turn, allowed to deposit ECM. Interestingly, ECM deposited by iMEFs cultured in conditioned medium from tRNA_{Met}-overexpressing cells had significantly enhanced ability to support cell migration (Figure 3B), indicating that tRNA_{Met} levels control the synthesis and secretion of soluble ECM components rather than the ability of fibroblasts to remodel these and incorporate them into insoluble ECM.

Cell migration may be influenced by the mechanical properties of the ECM. ECM stiffness promotes cell migration, and certain secreted enzymes (such as lysyl oxidases and transglutaminases) catalyze the formation of covalent cross-links between amino acid chains to increase ECM stiffness. We used atomic force microscopy (AFM) to measure the mechanical properties of ECM and found that cell-derived matrix generated in the presence of conditioned medium from tRNA_{Met}-overexpressing cells did not consistently differ in its thickness or stiffness from that generated in the presence of conditioned medium from control cells (Figure 3C). Taken together, these data indicate that increased tRNA_{Met} expression supports deposition of a pro-migratory ECM via release of secreted components that do not influence the mechanical properties of the ECM.

Increased tRNA_{Met} Levels Drive a Secretome that is Enriched in Type II Collagen

We used SILAC-based mass spectrometry to quantitatively determine the influence of tRNA_{Met} levels on the complement of soluble proteins secreted by fibroblasts. We labeled iMEF-tRNA_{Met} and iMEF-vector cells with light and heavy SILAC amino acids, respectively (forward [Fw] labeling), and with heavy and light amino acids, respectively (reverse [Rev] labeling). We then prepared cell extracts, and for the secretome, we used an affinity bead (Strataclean)-based approach to enrich for proteins secreted into serum-free conditioned medium [15]. Proteomes were detected using an LTQ-Orbitrap, and mass spectrometry data were analyzed with the MaxQuant computational platform, which identified proteins in the proteome and in the secretome (Tables S1 and S2) with high correlation between Fw and Rev data. Analysis of the cellular proteome indicated that increased tRNA_{Met} expression supports deposition of a pro-migratory ECM via release of secreted components that do not influence the mechanical properties of the ECM.

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Figure 2. Increased tRNA_{Met} Levels Drive a Secretome that is Enriched in Type II Collagen

(A) We used iMEF-tRNA_{Met} and iMEF-vector cells with light and heavy SILAC amino acids, respectively (forward [Fw] labeling), and with heavy and light amino acids, respectively (reverse [Rev] labeling). We then prepared cell extracts, and for the secretome, we used an affinity bead (Strataclean)-based approach to enrich for proteins secreted into serum-free conditioned medium [15]. Proteomes were detected using an LTQ-Orbitrap, and mass spectrometry data were analyzed with the MaxQuant computational platform, which identified proteins in the proteome and in the secretome (Tables S1 and S2) with high correlation between Fw and Rev data. Analysis of the cellular proteome indicated that increased tRNA_{Met} expression supports deposition of a pro-migratory ECM via release of secreted components that do not influence the mechanical properties of the ECM.

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In particular, type II collagen secretion by fibroblasts was upregulated by at least 50-fold following tRNA\textsubscript{Met} overexpression in these cells (Figures 4B and 4C; Table S2). Furthermore, specificity in the nature of the secreted ECM components was observed in iMEF-tRNA\textsubscript{Met} cells, which did not necessarily correlate with protein abundance (Figure 4D). For example, despite the observation that fibronectin was one of the most-abundant proteins in the secretome, its levels were not significantly altered following tRNA\textsubscript{Met} overexpression (Figures 4C and 4D). Upregulated expression of a given protein is commonly associated with increased recruitment of its mRNA(s) to actively translating ribosomes (polysomes). We used sucrose density gradients to determine whether increased tRNA\textsubscript{Met} levels were able to drive polysomal recruitment of collagen II mRNA. We found no indication that the monosome/polysome ratio differed between iMEF-vector and iMEF-tRNA\textsubscript{Met} cells (Figure S5A), which is consistent with our observations that the overall levels of cellular protein synthesis were unaffected by increased tRNA\textsubscript{Met} levels (Figures S2C and S3C). However, iMEF-tRNA\textsubscript{Met} cells displayed markedly increased polysomal content of collagen II mRNA (Figure S5B). Conversely, polysomal association of mRNAs for fibronectin and the ARPP0 ribosomal protein with polysomes was unaffected by increased tRNA\textsubscript{Met} levels (Figures S5C and S5D).

Immunostaining of ECM deposited by two independent tRNA\textsubscript{Met}-overexpressing cell lines confirmed that upregulation of soluble secreted type II collagen was also reflected by its increased incorporation into the insoluble ECM (Figure 4E). We examined whether levels of tRNA\textsubscript{Met} in the stroma influences the collagen II content of tumors in vivo. To do this, we introduced LLC cells into wild-type and 2+tRNA\textsubscript{Met} mice as syngeneic allografts, allowed the tumors to grow for 21 days, and then stained them for collagen II. Collagen II deposition was clearly visible in the spaces between tumor cells, and quantitative analysis indicated that the collagen II content of tumors grown in 2+tRNA\textsubscript{Met} animals was significantly higher than it was in tumors from littermate control animals (Figures 5A and 5B).

**Type II Collagen Secretion Is Required for tRNA\textsubscript{Met} to Drive Production of Pro-tumorigenic ECM**

To assess whether type II collagen secretion is responsible for the ability of tRNA\textsubscript{Met} to deposit a pro-migratory ECM, we generated cell-derived matrices from iMEF-tRNA\textsubscript{Met} cells transfected with type II collagen-specific siRNA or non-targeting control. Knockdown of type II collagen in iMEF-tRNA\textsubscript{Met} cells resulted in conditioned medium with decreased ability to enable iMEFs to generate ECM that supported fibroblast migration (Figures 6A and S6A–S6C). Furthermore, knockdown of type II collagen (with two independent siRNA sequences) in primary lung fibroblasts isolated from 2+tRNA\textsubscript{Met} mice similarly compromised the ability of the ECM derived from these cells to support increased migration of endothelial cells (Figure 6B).

ECM was also made from iMEFs in which type II collagen levels had been suppressed using CRISPR genome editing (MEF-tRNA\textsubscript{Met}/collagen II CRISPR; Figures S6D and S6E). Immunostaining indicated significantly decreased type II collagen.
incorporation into ECM deposited by MEF-tRNA\textsubscript{Met}/collagen II CRISPR cells (Figure 6C, left panel). Consistently, ECM generated by MEF-tRNA\textsubscript{Met}/collagen II CRISPR fibroblasts had decreased ability to support increased cell migration (Figures 6C, right panel, and S6F).

We then determined whether collagen synthesis was required for generation of a pro-tumorigenic microenvironment in vivo. To do this, we used ethyl-3,4-dihydroxybenzoate (DHB), a prolyl hydroxylase inhibitor that is well-established to compromise collagen processing in vivo and thus prevents collagen secretion in vivo [16]. Indeed, DHB administration significantly reduced the collagen II content of allografted tumors in both wild-type and tRNA\textsubscript{Met} mice (Figure 5A and 5B). However, whereas DHB treatment did not affect vascularity or growth of LLC subcutaneous allografts in WT mice, it significantly opposed the increased vascularity and growth of tumors seen in 2+tRNA\textsubscript{Met} mice (Figure 5C), indicating that collagen production is indeed required for tRNA\textsubscript{Met} to drive tumor growth and angiogenesis in vivo.

**Type II Collagen Expression Predicts Poor Prognosis in Ovarian Cancer**

To determine whether the relationship between increased collagen II levels and tumor progression is observable in human cancer, we stained a tissue microarray containing 62 cases of primary ovarian tumors. The decision to focus on this cancer type was influenced by reports that ovarian cancer aggressiveness is particularly associated with increased angiogenesis [17]. In univariate analysis, high collagen II histoscores were associated with significantly decreased survival following resection of primary ovarian tumors (Figure 7), and this was particularly apparent in high-grade serous carcinoma (Figure S7). Indeed, patients with tumors displaying a collagen II histoscore greater than 50 have a 2.992-fold increased hazard ratio for death (p = 0.0002). These data indicate that collagen II expression is a significant factor in driving the progression of ovarian cancer.

**DISCUSSION**

Regulation of the cellular tRNA repertoire is thought to dictate selectivity of translational programs to influence cancer progression and to define cell fates [2]. Indeed, the tRNA repertoire...
differs between proliferating and differentiated cells, and this is thought to influence the translational landscape according to codon usage [2, 18]. Thus, an mRNA with a preponderance of a given codon will be translated more rapidly if the tRNA recognizing that codon is abundant. Consequently, one might anticipate that increased tRNA^\text{Met}\ (\text{which recognizes AUG at translation start sites}) would generally increase translation initiation and globally upregulate protein synthesis and cell growth [6].

However, our transgenic approach has given no indications that increased tRNA^\text{Met} levels promote cell growth or proliferation in a cell-autonomous fashion, nor does this greatly influence the quantity or profile of cellular protein synthesis. Instead, we have found that increased levels of tRNA^\text{Met} lead to highly selective adjustments to the protein synthesis landscape that are restricted to certain secreted proteins—particularly the ECM component, type II collagen. There is evidence from studies in Drosophila that elevated tRNA^\text{Met} controls production of secreted proteins. Introduction of one extra copy of the tRNA^\text{Met} gene into flies is sufficient to markedly enhance larval growth [7].

Consistent with our findings, this is primarily not attributable to a cell-autonomous drive to cell growth and proliferation in the developing organs. Rather, growth is driven by an endocrine relay in which increased tRNA^\text{Met} in fat body cells promotes secretion of factor(s) that trigger release of insulin-like peptides from brain neurosecretory cells that, in turn, circulate throughout the larvae to promote growth of developing organs and tissues [7].

ECM production constitutes a significant commitment to the cell (both in terms of load on the ER and the biosynthetic machinery), indicating the need for special mechanisms to be in place to control the translation of ECM proteins and other abundant components of the fibroblast secretome. Expression of type I collagens is known to be controlled at the level of mRNA stability and translation, due to unique 5′ stem loops in their 5′ UTRs [19], and the production of these fibrillar collagens by the stroma is established to influence cancer growth and metastasis [20, 21].

Type II collagen is also a fibrillar collagen; however, little is known about how its production might be controlled post-transcriptionally. Although the sensitivity of collagen II production to tRNA^\text{Met} levels indicates that translation initiation is likely a key point of...
Figure 5. Collagen II Production Is Required for tRNA\textsubscript{Met} to Drive Production of a Pro-tumorigenic ECM

1 × 10\textsuperscript{6} LLC cells were injected subcutaneously into either WT or 2+tRNA\textsubscript{Met} transgenic male mice, followed by daily administration of ethyl-3,4-dihydroxybenzoate (DHB) (40 mg/kg) or vehicle control. Tumors were harvested 21 days following injection. Tumor volumes were measured and tumor sections were stained for collagen II or endomucin.

(A and B) Collagen II staining in LLC tumors harvested from WT and 2+tRNA\textsubscript{Met} mice treated with vehicle or DHB as indicated (A). Collagen II staining was quantified by using the Image J “threshold color” plug-in to highlight the brown staining in the left panels and to render this as a grayscale image as shown in the graph in (B) (legend continued on next page).

(A) WT - vehicle
(B) WT - DHB

(C) Vessels per mm\textsuperscript{2}
(D) Tumor volume (mm\textsuperscript{3})

(A) 2+tRNA\textsubscript{Met} - vehicle
(B) 2+tRNA\textsubscript{Met} - DHB

(legend continued on next page)
control of type II collagen synthesis, there are indications that the mechanism for this might not be straightforward. Indeed, in some cells (including those used for the polysome analysis displayed in Figure S5), we find that tRNA^{Met} overexpression increases the cellular levels of mRNA for type II collagen (not shown), indicating that tRNA^{Met} levels may influence transcription or stability of mRNAs for ECM proteins. Thus, despite clear indications that tRNA^{Met} levels selectively control ECM protein synthesis in a way that influences tumor outcomes, further work will be needed to fully establish the molecular mechanisms through which tRNA^{Met} levels influence type II collagen synthesis and the contribution made by the translational machinery to this non-cell-autonomous drive to tumor progression.

Despite the presence of a number of tRNA^{Met}-sensitive components in the fibroblast secretome, it is clear that suppression of just one of these, type II collagen, is sufficient to negate the tRNA^{Met} phenotype. Moreover, we have shown that stromal levels of type II collagen are strongly associated with aggressiveness of serous ovarian carcinoma, indicating the importance of this tRNA^{Met}-sensitive secreted factor to the progression of a human cancer type. However, we have been unable to quantitatively determine tRNA^{Met} levels in human tumor microarrays, and it will be necessary to do this to conclusively determine whether stromal tRNA^{Met} drives type II collagen synthesis in human tumors in the same way as we have demonstrated by performing allografts in our 2+tRNA^{Met} mouse model. Type II collagen is an abundant component of cartilage and is produced in large quantities by chondrocytes. By contrast, normal fibroblasts are not thought to secrete collagen II, and consistently, our SILAC data show that this collagen is a relatively minor component of the fibroblast secretome. Thus, acquisition of type II collagen secretion when tRNA^{Met} levels are increased may represent a shift to a more-chondrocyte-like phenotype, and this may be consistent with the role for the tRNA repertoire in influencing differentiation state [2]. In view of this, one might anticipate that increased tRNA^{Met} levels may be associated with production of ECM with cartilaginous features, such as increased stiffness. However, our AFM data show clearly that ECM produced by fibroblasts with increased tRNA^{Met} levels does not have consistently altered compliance, indicating that type II collagen content is not necessarily linked to stiffness. In addition to providing mechanical support, a key role of the ECM is to act as a reservoir for growth factors and other bioactive proteins. The tRNA^{Met} secretome contains a number of pro-tumorigenic growth factors (such as frizzled-related protein and TGF-β2), and we are currently considering the possibility that type II collagen production is associated with secretion of these growth factors and their incorporation into the ECM, increasing their presentation to tumor and endothelial cells.
Targeting the translational initiation machinery with drugs has been shown to have sufficiently selective effects on protein synthesis in tumor cells to influence cancer progression. Indeed, eIF4A, which is the molecular target for the promising anticancer agent silvestrol, favors translation of key oncoproteins such as Myc, Notch, and BCL2 [22]. Our findings indicate that control of tRNA<sub>Met</sub> levels in stromal fibroblasts is key to the generation of a collagen-II-rich pro-tumorigenic microenvironment, and this will provide further impetus to the search for drugs to manipulate translation initiation and ECM synthesis in cancer.

Our data also have implications for understanding the mechanisms of tumor angiogenesis. Although stromal fibroblasts are known to contribute to angiogenesis in tumors, the mechanisms through which fibroblasts promote tumor angiogenesis are incompletely understood [23]. The data presented here suggest that the secretion of a pro-migratory ECM by stromal fibroblasts is an important mechanism via which stromal fibroblasts promote angiogenesis in cancer.

EXPERIMENTAL PROCEDURES

Cell Culture

Primary MEFs were isolated from embryonic day 13 (E13)–E15 WT or 2+tRNA<sub>Met</sub> embryos. After sterile dissection from the yolk sac and removal of the fetal liver and head, the embryo was disaggregated by forceful pipetting and cells pelleted by centrifugation and resuspended in DMEM supplemented with 10% FBS, 200 μM L-glutamine, 100 U/ml penicillin streptomycin, 0.25 μg/ml Fungizone, and 50 μM β-mercaptoethanol. Cells were cultured on 0.1% gelatin-coated plates and used as described in the text. For routine culture, human umbilical vein endothelial cells (HUVECs) from pooled donors (TCS Cell Works) were seeded onto 0.1% gelatin-coated plates and cultured in M199 medium supplemented with 20% FBS, 0.1 mg/ml bovine brain endothelial mitogen, 1 ng/ml heparin, and 100 U/ml penicillin streptomycin. HUVECs were used for experiments at passages 4–6. Both B16F0 and LLC cells were cultured in DMEM supplemented with 10% FBS.

Animal Experimentation

All animal work was carried out with ethical approval (University of Glasgow and Institute of Cancer Research) in dedicated barrier animal facilities proactive in environmental enrichment and in accordance with the revised Animals (Scientific Procedures) Act 1986 and the EU Directive 2010/63/EU.

2+tRNA<sub>Met</sub> Mice

Two copies of the murine tRNA<sub>Met</sub> gene (tRNA<sub>78</sub>) including ~140 bp of flanking sequence, were targeted into the HPRT locus of HM1 embryonic stem cells and used to generate chimeric mice. 2+tRNA<sub>Met</sub> transgenic offspring were derived by crosses to C57Bl/6 mice and maintained on a C57Bl6/J background.

Syngeneic Allografts

Cells were injected subcutaneously into the right flank of either WT or 2+tRNA<sub>Met</sub> transgenic mice. Tumors were harvested at 21 or 22 days post-implantation (for LLC and B16) or at endpoint (for melanocytes), and final tumor volume was calculated using the formula (L x W x W)/2, where L is the longest dimension and W is the shortest dimension. For blood vessel identification, tumors were fixed and stained for endomucin. Vessels were counted across the entire tumor section and are reported as vessels/mm<sup>2</sup>. Only tumors of a similar size were counted, with extremely large and extremely small tumors excluded to allow for reasonable biological comparison. Selection of tumors to analyze was made before blood vessel counts. Forty milligrams per kilogram DHB was administered daily by intraperitoneal injection.

Expression Vectors

Fragments containing tRNA<sub>Met</sub> were initially cloned into the multiple cloning site of the pLPCX vector using EcoRI. The pLPCX-tRNA<sub>Met</sub> construct and pLHCX final destination vector were digested using BgII and ClaI and ligated to produce the pLHCX-tRNA<sub>Met</sub>-containing vector. The vector was digested with HpaI and SnaBI and then re-ligated. For CRISPR gene editing, a tissue microarray (TMA) containing 62 cases of ovarian cancer was stained for type II collagen. Examples of tumor cores displaying low (left side) and high (right side) histoscores for collagen II are displayed in (A). (B) shows a Kaplan-Meier analysis of patient survival when the cohort is divided into tumors with low (histoscore < 50) and high (histoscore > 50) collagen II expression. Patients with tumors displaying a collagen II histoscore greater than 50 have a 2.992-fold increased hazard ratio for death; p = 0.0002 (log rank test); n = 62. See Figure S7 for Kaplan-Meier analysis of the relationship between collagen II levels in the high-grade serous subset of these tumors.

Figure 7. Type II Collagen Expression Predicts Poor Prognosis in Ovarian Cancer

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single-stranded oligonucleotides were designed to exon 1 of collagen II, annealed to make double-stranded oligos, and ligated into the linearized GeneArt CRISPR Nuclease Vector as per manufacturer’s instructions (Life Technologies).

qRT-PCR
RNA was prepared using Trizol (Life Technologies) and cDNA prepared using Quantitech Reverse Transcription Kit (QIAGEN), both as per manufacturer’s instructions, using 5, 10, and 20 ng RNA templates to ensure the reaction was quantitative. qPCR was conducted using SyBr green (Quanta) and the Bio-Rad CFX platform, with a 60°C annealing temperature and the primer pairs that are tabulated in the Supplemental Experimental Procedures.

siRNA Transfection
iMEFs were transfected with 10 nM collagen II siRNA (L-043139-01-0005 mouse Col2a1 SMARTpool; Dharmacon), or non-targeting siRNA, using the Amaxa Nucleofector system, solution R, program T-20, as per manufacturer’s instructions (Lonza). Cells were used for experiments at 48 hr post-transfection.

ECM Generation and Cell Migration
The preparation of cell-derived matrix was performed as previously described [14], as was time-lapse analysis of cell migration [13].

AFM
To measure the thickness of ECMs following wound scratch, AFM force spectroscopy and contact imaging were performed using a JPK NanoWizard II Bio AFM in combination with a Bruker MLCT cantilever. A Nanoworld Arrow TL-1 cantilever with bead attached was used for force spectroscopy.

SILAC Mass Spectrometry
iMEF-vector and iMEF-tRNAiMet cells were labeled with light (Arg0/Lys0; Sigma) and heavy amino acids (Arg10/Lys8; Cambridge Isotopes Laboratories). In the case of the forward experiment, iMEF-vector cells were labeled with light amino acids and iMEF-tRNAiMet cells with heavy amino acids, whereas in the reverse experiment, iMEF-vector cells were labeled with heavy amino acids and iMEF-tRNAiMet cells with light amino acids. Briefly, cells were passaged in SILAC DMEM containing light or heavy amino acids supplemented with 10% FBS for three passages and then transferred to media containing light or heavy amino acids supplemented with 10% dialyzed FBS. Cells were cultured for 48 hr at 37°C/5% CO₂, washed twice with PBS, and transferred into serum-free media for 6 hr at 37°C/5% CO₂. The media was then acidified to pH 5 with 10% TFA and 10 μl Strachatein beads added per 1 ml media. Beads were collected by brief centrifugation and proteins separated by electrophoresis. Proteins were digested with 0.05 μg/ml trypsin in 50 mM ammonium bicarbonate overnight. Extracted digests were stage tipped and analyzed on a linear trap quadrupole (LTQ)-Orbitrap Elite. Data were searched and quantified against Uniprot MOUSE using MaxQuant [24].

Tissue Microarray and Immunohistochemistry
The SaPPrOC trial (Clinicaltrials.gov reference NCT01196741) was a randomised phase II study of weekly paclitaxel ± oral saracatinib in women with breast cancer (BCT). Tissue Microarray and Immunohistochemistry

REFERENCES


