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## The role of the *MTA* family and their encoded proteins in human cancers: molecular functions and clinical implications

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**Abstract** *MTA* (metastasis-associated gene) is a newly discovered family of cancer progression-related genes and their encoded products. *MTA1*, the first gene found in this family, has been repeatedly reported to be overexpressed along with its protein product MTA1 in a wide range of human cancers. In addition, the expression of *MTA1*/MTA1 correlates with the clinicopathological properties (malignant properties) of human cancers. MTA proteins are transcriptional co-repressors that function in histone deacetylation and are involved in the NuRD complex, which contains nucleosome remodeling and histone deacetylating molecules. MTA1 expression correlates with tumor formation in the mammary gland. In addition, MTA1 converts breast cancer cells to a more aggressive phenotype by repression of the estrogen receptor (ER)  $\alpha$  trans-activation function through deacetylation of the chromatin in the ER-responsive element of ER-responsive genes. Furthermore, MTA1 plays an essential role in c-MYC-mediated cell transformation. Another member of this family, MTA3, is induced by estrogen and represses the expression of the transcriptional repressor Snail, a master regulator of “epithelial to mesenchymal transitions”, resulting in the expression of the cell adhesion molecule E-cadherin and maintenance of a differentiated, normal epithelial phenotype in breast cells. In addition, tumor suppressor p53 protein is deacetylated and inactivated by both MTA1 and MTA2, leading to inhibition

of growth arrest and apoptosis. Moreover, a hypoxia-inducible factor-1 $\alpha$  (HIF-1 $\alpha$ ) is also deacetylated and stabilized by MTA1, resulting in angiogenesis. Thus, MTA proteins, especially MTA1, represent a possible set of master co-regulatory molecules involved in the carcinogenesis and progression of various malignant tumors. MTA proteins are proposed to be important new tools for clinical application in cancer diagnosis and treatment.

**Keywords** Metastasis-associated gene 1(MTA1) · Chromatin remodeling · Histone deacetylation · Gene expression · Protein modification · Cancer progression · Metastasis

### Abbreviations

MTA	Metastasis-associated gene/protein
HDAC	Histone deacetylase
NuRD	Nucleosome remodeling and histone deacetylation
ER	Estrogen receptor
HIF	Hypoxia-inducible factor

### Introduction

Recent advances in molecular biology have resulted in the discovery of a wide variety of new molecules involved in carcinogenesis and cancer progression. Although additional molecules related to cancer will be identified in the future, the existing and new molecules must fulfill two major requirements in order to be clinically useful as molecular targets for the diagnosis and treatment of human cancers. The first is that abnormalities in expression or structure of molecules of interest and their clinical relevance must be definitely demonstrated in human cancers by

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independent studies. The second is the underlying molecular mechanisms necessary for the molecules to exert their functions in carcinogenesis or cancer progression must be determined.

Among a number of cancer-related genes and molecules that have been discovered in the last few years, we identified a candidate metastasis-associated gene by use of a differential cDNA screening method. Thus, we identified a gene that was abundantly overexpressed in highly metastatic rat mammary adenocarcinoma cell lines compared to poorly metastatic cell lines [1, 2]. When this gene was sequenced, it was revealed as a completely novel gene without any homologous or related genes in the database. The rat gene was named *mtal* (metastasis-associated gene 1). A homologous gene was also expressed in human cancer cell lines [1], and its human cDNA counterpart, *MTA1*, was cloned by our group in 2000 [3]. Using surgically resected human tissues, we showed that high levels of *MTA1* mRNA expression were clinicopathologically correlated to the invasive and growth properties of gastrointestinal cancers, including esophageal, gastric and colorectal cancers [4, 5]. Subsequently, several reports from independent research groups followed our observations and showed similar correlations between *MTA1* expression and the malignant potentials of human cancers.

Several genes related to *MTA1* have now been identified, indicating *MTA1* consists of a gene family, which we now call the “*MTA* family”. Further studies on molecular biological and biochemical properties of the *MTA* family have shown that the gene products of the main members of the family (*MTA1*, *MTA2*, and *MTA3*) are tightly associated in a protein complex called NuRD (nucleosome remodeling and histone deacetylation), which has transcriptional regulatory function via histone deacetylation and chromatin remodeling. At the moment, the *MTA* family has attracted widespread attention as one of the key molecules that plays an indispensable role in the genesis and progression of a wide variety of cancers [6–8]. In this brief review, we will examine the significance of the expression of *MTA* family members in human cancers and the important molecular mechanisms that are currently known by which *MTA* proteins exert their functions. Finally, future directions for clinical applications of this protein family for the diagnosis and treatment of human cancers will be discussed.

### Members of the *MTA* family and their protein structures

At present, the *MTA* proteins represent a family of gene products encoded by three distinct genes (*MTA1*, *MTA2*, and *MTA3*) and six reported isoforms (*MTA1*, *MTA1s*, *MTA1-ZG29p*, *MTA2*, *MTA3*, and *MTA3L*). The molecular masses

of the gene products of *MTA1*, *MTA2*, and *MTA3* are approximately 80, 70, and 65 kDa, respectively. The nucleotide and protein alignment homologies and the phylogenetic comparative analyses are discussed elsewhere [8, 9].

Except for ZG-29p, the *MTA* family sequences contain several common domain structures [10]. One of these, the BAH (bromo-adjacent homology) domain is involved in protein–protein interactions. Another, the SANT (SWI, ADA2, N-CoR, TFIIB-B) domain shares a high degree of homology with the DNA-binding domain of the Myb-related proteins, suggesting that this domain may be involved in DNA-binding. The ELM (egl-27 and *MTA1* homology) domain has an unknown function [11]. *MTA* family members also contain a highly conserved GATA-type zinc finger motif, which indicates a direct interaction with DNA [3]. *MTA1* has two src-homology (SH)-binding motifs at its C-terminal region, which are known to be important in signal transduction involving many kinase, adaptor and scaffolding proteins [1, 10]. Similar SH2- and SH3-binding domains are also found in *MTA2* and *MTA3*. These common domain structures clearly show that the *MTA* family is involved in protein–protein and DNA-binding interactions, indicating possible functions in signal transduction and transcriptional regulation.

*MTA* proteins contain basic nuclear localization signals [1, 10]. They also localize in the nucleus in many cancer cells [4, 8]. However, *MTA1* localizes to both the cytoplasm and nucleus in some tumors [12–14]. *MTA3* also localizes to the nucleus, but it has no apparent nuclear localization signal [15]. *MTA1s*, a short splice-variant of *MTA1*, is predominantly localized in the cytoplasm [16].

### The expression of *MTA* proteins in various cancers and its clinicopathological and biological relevance

#### Clinicopathological relevance of the increased *MTA1* expression in human cancer tissues

Since the first report by us showing that the up-regulation of *MTA1* expression was significantly correlated to the malignant properties of human gastric and colorectal cancers [4], many researchers have been investigating the expression levels of *MTA* family members, especially *MTA1*, in various human cancers. This has revealed that the expression levels of *MTA* family members have clinicopathological significance (The data are summarized in Table 1).

#### Breast cancer

*MTA1* was identified as a candidate progression molecule that was associated with breast cancer metastasis [1, 2] and

growth (the antisense RNA of MTA1 inhibited the growth of highly metastatic breast cancer cell lines [3]). The involvement of MTA1 in the carcinogenesis or progression of human breast cancer was also shown by other data using clinical samples. For example, Martin et al. [17] mapped the

chromosomal locus 14q that might be responsible for axillary lymph node metastasis in human breast cancers by comparing the rate of loss of heterozygosity between node-positive and -negative breast cancers. They found that the *MTA1* gene was contained in that gene locus, suggesting that

**Table 1** Clinicopathological implications of the increased MTA1 expression in various human cancer tissues

Type of cancer	Method	Clinicopathological implications	Reference
Breast cancer	LOH	Higher LN meta.	[17]
	IHC	Earlier recurrence	[18]
	IHC	Higher tumor grade Higher MVD (angiogenesis)	[19]
<i>Gastrointestinal cancer</i>			
Esophageal	RT-PCR	Deeper adventitial invasion Higher LN meta.	[5]
	IHC	Deeper adventitial invasion Higher LN meta. More advanced stage Poorer prognosis	[21]
Gastric	RT-PCR	Deeper serosal invasion Higher LN meta.	[4]
Colorectal	RT-PCR	Deeper wall invasion Higher LN meta.	[4]
	RT-PCR	Higher expression in cancer tissue	[20]
<i>Carcinoid</i>			
Gastric	RT-PCR	Deeper tumor invasion	[25]
Small intestine	RT-PCR	Malignant carcinoid More liver and LN meta.	[22]
Appendiceal	RT-PCR	Significant increase in malignant tumors	[26]
Pancreatic	IHC	Poorer prognosis (in combination with HDAC1)	[28]
Hepatocellular	RT-PCR	Shorter disease-free survival	[29]
	IHC	Larger tumor size More vascular invasion	[12]
	IHC	More microvascular invasion Higher recurrence rate Poorer survival	[30]
<i>Other cancers</i>			
NSCLC	RT-PCR	Larger tumor size Higher LN meta.	[31]
Thymoma	RT-PCR	More advanced stage	[33]
Ovarian	RT-PCR	Higher LN meta.	[32]
	IHC	More advanced stage Higher FIGO staging	[34]
Prostate	IHC	Metastatic prostate ca.	[35]
Lymphoma	Microarray	Highest expression in diffuse B-cell lymphoma	[36]
HNSCC	Microarray	Higher LN meta.	[37]
	IHC	Higher LN meta. More advanced stage Deeper wall invasion	[38]

NSCLC non-small cell lung cancer, HNSCC head and neck squamous cell carcinoma, IHC immunohistochemistry, LOH loss of heterozygosity, RT-PCR reverse transcription-polymerase chain reaction, MVD microvessel density, LN meta. lymph node metastasis

*MTA1* is a strong candidate for a breast cancer metastasis-promoting gene. Furthermore, using immunohistochemistry they examined the *MTA1* protein expression in primary human breast cancer samples and demonstrated that node-negative breast cancers with overexpression of *MTA1* protein had a higher risk of disease relapse similar to node-positive tumors. Thus, the overexpression of *MTA1* may be a useful predictor of early disease relapse [18].

Jang et al. [19] also showed that *MTA1* overexpression was closely associated with higher tumor grade and high intratumoral microvessel density in surgically resected human breast cancers, suggesting that *MTA1* may be a useful predictor of an aggressive phenotype and a possible angiogenesis-promoting molecule in breast cancer.

### Gastrointestinal cancer

By using a reverse-transcription polymerase chain reaction (RT-PCR) method, we demonstrated that the higher expression of *MTA1* mRNA in surgically resected human gastric and colorectal cancer specimens compared to the paired normal counterpart tissues was significantly correlated to the depth of cancer invasion and lymph node metastasis [4]. This study was the first to demonstrate the clinical relevance of *MTA1* expression to the malignant potentials of human cancers. Higher expression of *MTA1* mRNA was also shown in colorectal cancers compared to the normal counterpart tissue by another group [20].

Using a RT-PCR method, we found that human esophageal squamous cell cancers overexpressed *MTA1* mRNA. The overexpressing cancer cells showed significantly higher frequencies of adventitial invasion and lymph node metastasis and tended to have a higher rate of lymphatic involvement [5]. Using immunohistochemistry, we further examined the protein expression level of *MTA1* in human esophageal squamous cell cancers and reconfirmed the results obtained by RT-PCR [21]. In this study, we also demonstrated that *MTA1* was a predictor of poor prognosis after surgery [21].

In another observation, Kidd et al. [22] showed that it was useful to examine the expression of *MTA1* mRNA and *MTA1* protein in order to determine the malignant potential and the propensity to metastasize of small intestinal carcinoid (enterochromaffin cell) tumors. When compared to nonmetastatic primary tumors, the expression of *MTA1* was increased in malignant small intestinal carcinoids and in metastases to liver and lymph nodes [22–24]. This same group further reported that *MTA1* was a good candidate genetic molecular marker to discriminate gastric carcinoids from other gastric neoplasms [25] as well as malignant appendiceal carcinoids from benign tissue [26]. In these studies, *MTA1* was thought to be a good marker to define the malignancy of carcinoid tumors.

In addition to cancers of the gastrointestinal tract, the involvement of *MTA1* overexpression in carcinogenesis and cancer progression was shown in other gastrointestinal tumors, such as pancreatic cancers and hepatocellular carcinomas. Iguchi et al. [27] examined *MTA1* mRNA expression in pancreatic cancer cell lines and resected pancreatic cancer tissues and found that increased levels of *MTA1* mRNA expression might be involved in the progression of pancreatic cancer. Recently, Miyake et al. [28] showed the expression level of *MTA1* protein correlated with poorer prognosis of pancreatic cancer patients.

The possible association of *MTA1* expression with the malignant properties of hepatocellular carcinomas (HCC) was first reported by Hamatsu et al. [29]. In this study, *MTA1* mRNA level was assessed by RT-PCR in resected human HCC tissues, and its high expression predicted a lower disease-free survival rate after curative hepatectomy for HCC. Using immunohistochemistry, Moon et al. [12] examined *MTA1* protein expression in resected human HCC specimens. They showed that overexpression of *MTA1* was associated with HCC growth and vascular invasion and that nuclear localization of estrogen receptor (ER)  $\alpha$  inversely correlated with *MTA1* expression, suggesting that *MTA1* was involved in negative regulatory mechanisms. Ryu et al. [30] reported that *MTA1* was closely associated with microvascular invasion, frequent postoperative recurrence, and poor prognosis in patients with HCC, especially in those with hepatitis B virus (HBV)-associated HCC.

### Other cancers

The relationship between *MTA1* expression and malignant properties, such as invasion and metastasis, has been investigated in many other carcinomas and sarcomas. High expression of *MTA1* mRNA was correlated clinicopathologically with lymph node metastasis of human non-small cell lung cancers [31] and ovarian cancers [32], and to the advanced stage and invasiveness of thymomas [33]. Dannenmann et al. [34] reported that overexpression of *MTA1* protein in ovarian cancer was significantly correlated to more advanced stage and higher FIGO staging. The potential role of *MTA1* protein expression has also been suggested in the progression of human endometrial carcinomas [14]. In prostate cancers, Hofer et al. [35] showed that metastatic prostate tumors demonstrated significantly higher intensities of *MTA1* protein expression and higher percentages of tissue cores staining positive for *MTA1* than in clinically localized prostate cancers or benign prostate tissues. The high expression of *MTA1* in diffuse B-cell lymphomas was also reported in human cases [36].

Using DNA microarray analysis, Roepman et al. [37] investigated gene expression patterns in lymph node

metastases of head and neck squamous cell carcinomas. They showed that the *MTA1* gene was the only single gene that showed consistently changed expression between numbers of matched pairs of primary tumor and lymph node metastases. Recently, further evidence was reported showing that overexpression of MTA1 protein in oral squamous cell carcinoma correlated to higher lymph node metastasis, deeper wall invasion and more advanced stage [38].

#### Biological relevance of MTA proteins to carcinogenesis and cancer progression

In addition to the clinicopathological evidences mentioned above, the biological relevance of MTA proteins to carcinogenesis and cancer progression has been made much clearer by the following important experiments.

The direct evidence to show the association of MTA1 expression with breast cancer malignant properties was first obtained by Mazumdar et al. in 2001 [39]. They demonstrated that forced expression of the MTA1 protein in breast cancer cell line MCF-7 was accompanied by enhancement of the ability of cells to invade an artificial matrix and to grow in an anchorage-independent manner. They also showed that the enhancement was associated with the interaction between MTA1 protein and histone deacetylase, resulting in a repression of ER $\alpha$ -mediated transcription (This will be discussed in more detail later).

The above study was extended by further experiments by the same group where they showed direct in vivo evidence of the involvement of MTA1 in the carcinogenesis of breast cancer in an animal model [10, 40]. This group established transgenic mice that overexpressed MTA1 protein. The MTA1-transgenic mice showed an inappropriate development of mammary glands, and the mice eventually developed hyperplastic nodules and mammary tumors, including adenocarcinomas. Most interestingly, MTA1-transgenic mice were accompanied by high incidence of spontaneous B cell lymphomas, including diffuse large B cell lymphomas [13, 41].

The clinicopathological correlation of MTA1 overexpression with squamous cell carcinomas was reinforced by the experimental results of Mahoney et al. [42]. They transfected *MTA1* cDNA into immortalized human keratinocytes and clearly showed that forced expression of *MTA1* contributed to several aspects of enhanced metastatic behavior, including increased migration, invasion and survival in the anchorage independent state of the immortalized keratinocytes. Furthermore, Qian et al. [43] inhibited *MTA1* expression by RNA interference (RNAi) in a human esophageal squamous cell carcinoma cell line and showed the significant inhibition of in vitro invasion and migration properties of the cancer cells.

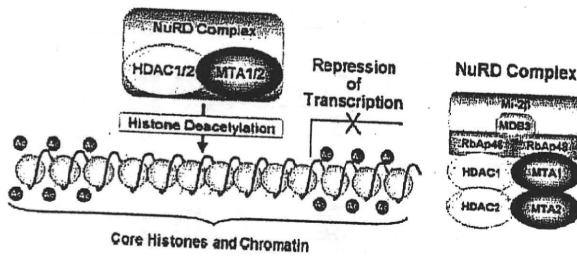
Direct evidence showing the role of MTA1 in the progression of pancreatic cancer was provided by Hofer et al. [44]. They transfected *MTA1* cDNA into the pancreatic cell line PANC-1 and demonstrated that enhanced expression of MTA1 promoted the acquisition of an invasive and metastatic phenotype and that it enhanced the malignant potentials of pancreatic adenocarcinomas by modulation of the cytoskeleton via IQGAP1.

#### Molecular mechanisms of the MTA family, especially MTA1, in carcinogenesis and cancer progression

As mentioned above, it was demonstrated by different approaches and by different laboratories that MTA1 overexpression was closely correlated with carcinogenesis and cancer progression of a wide range of cancers originating in disparate organs and tissues. This strongly indicates that MTA1 may be one of the important key molecules in the cancer progression field. Thus, it will be absolutely necessary to clarify the molecular mechanisms in which MTA family members exert their functions for the clinical utilization of MTA proteins for diagnosis or treatment of human cancers. Here, we introduce the several important functions of MTA proteins that have been clarified, especially those that are concerned with carcinogenesis and cancer progression.

#### Nucleosome remodeling and histone deacetylation complex and transcriptional regulation

The first notion about the molecular and biochemical functions of MTA1 was accidentally obtained by four independent groups in 1998–1999 [9, 45–48]. In these studies, two disparate chromatin modifying activities, ATP-dependent nucleosome remodeling activity and histone deacetylation, were functionally and physically linked in the same protein complex. This complex has been named the NuRD (Nucleosome remodeling and histone deacetylation), and it contains histone deacetylase (HDAC) 1, HDAC2, the histone binding proteins RbAp46/48 and the dermatomyositis-specific autoantigen Mi-2, which has been shown to have transcription repressing activity. Xue et al. [46] reported that MTA1 protein was found in the NuRD complex, and it had strong transcription repressing activity. Subsequently, Zhang et al. [47] reported that a protein similar to MTA1 (named MTA2) was also a component of the NuRD complex and that MTA2 is highly expressed in rapidly dividing cells. Later, MTA3 was identified as an estrogen-inducible gene product that forms a distinct NuRD complex [15]. We also reported the physical interaction between MTA1 and HDAC1 [49] (Fig. 1).



**Fig. 1** The fundamental function of MTA proteins. The fundamental function of MTA proteins is chromatin remodeling and histone deacetylation, resulting in repression of transcription. MTA proteins are included in the protein complex named NuRD, which also contains histone deacetylases (HDAC1 and 2), major DNA binding protein 3 (MDB3), histone binding proteins RbAp46/48 and the dermatomyositis-specific autoantigen Mi-2, and has strong transcription repressing activities

Thus, the fundamental functions of the MTA family members appear to be exerted through a NuRD complex that has chromatin remodeling and histone deacetylating properties (There is also deacetylating property of non-histone proteins in the NuRD complex). In addition, the MTA-NuRD complex shows transcriptional repression activities [6–8, 10, 50]. Although all MTA family proteins are found in NuRD complexes, these proteins form distinct NuRD complexes that are thought to target different sets of promoters [9].

#### Repression of the transactivating function of ER $\alpha$ by MTA proteins

Although the involvement of MTA proteins in NuRD complexes suggested that such complexes might function in chromatin remodeling and histone deacetylation, a direct target of MTA proteins was first identified by Mazumdar et al. in 2000 [39]. MTA1 was identified as a molecule induced by a growth factor, heregulin-beta1 (HRG), which is a natural ligand of the human epidermal growth factor receptors HER3 and HER4 that can also transactivate HER2 (c-erbB-2) in human breast cancer cell lines. They showed that MTA1 directly interacted with the ligand-binding domain of ER $\alpha$  and that HRG stimulated the association of MTA1 and HDAC2 on the chromatin of an ER-responsive element (ERE) in the promoters of the estrogen responsive genes, such as pS2 and c-myc. This explains the phenomenon that activation of HRG/HER2 pathway in ER-positive breast cancers results in the suppression of ER $\alpha$  functions, resulting in more invasive and aggressive phenotypes observed in ER-negative breast cancers [51]. The repressive function of MTA1 on ER $\alpha$  is mediated through histone deacetylation by HDAC1 and HDAC2, suggesting that MTA1 has a potent corepressor function on the transactivation function of ER $\alpha$  through histone deacetylation (Fig. 2a). MTA2 has also been shown

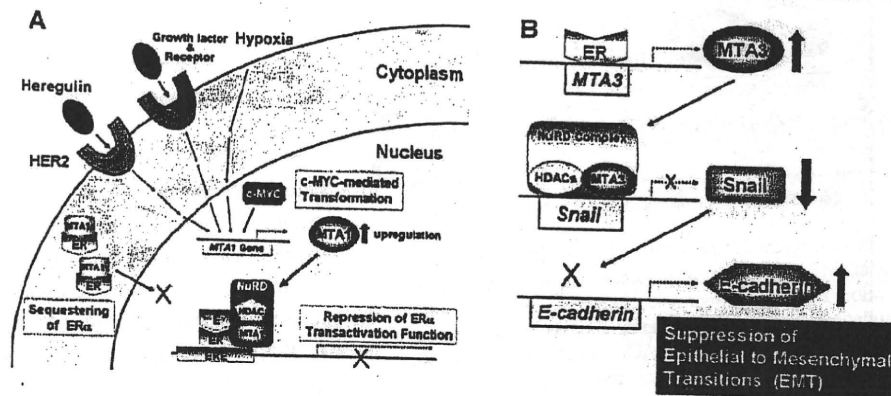
to physically interact with ER $\alpha$  and to repress its transactivating function. Furthermore, overexpression of MTA2 rendered cells unresponsive to estrogen and suppressed estrogen-induced colony formation in breast cancer cells [52] (Fig. 2a, b).

Recently, Khaleque et al. [53] showed that MTA1 binds to a heat shock factor 1 (HSF1), the transcriptional activator of the heat shock genes, in vitro and in human breast carcinoma samples. They demonstrated that HSF1-MTA1 complex formation was strongly induced by HRG and that the complex was incorporated into the NuRD complex and participated in repression of estrogen-dependent transcription in breast cancer cell treated with HRG.

Following the report by Mazumdar et al. [39], the same research group reported that several molecules, such as ménages a trios 1 (MAT1), MTA1-interacting coactivator (MICoA) and nuclear receptor interacting factor 3 (NRIF3), all interact with MTA1 and repress the transactivation function of ER $\alpha$  [8]. These three MTA1-binding proteins themselves have coactivator properties upon ER $\alpha$  transactivation. Talukder et al. [54] identified MAT1, an assembly and targeting ring finger factor for cyclin-dependent kinase-activating kinase (CAK), as a MTA1-binding protein. The interactions between CAK and MTA1 apparently regulate the transactivation activity of ER $\alpha$  in a CAK-dependent manner in breast cancer cells. In contrast, MICoA-mediated ER $\alpha$  transactivation functions are opposed by MTA1 through the recruitment of HDACs [55]. Furthermore, the interactions between MTA1 and NRIF3, an estrogen-inducible gene, may be important in modulating the sensitivity of breast cancer cells to estrogen [56]. Singh et al. [57] identified another MTA1-binding partner, Lim-only protein 4 (LMO4). LMO4 was found to be a component of the MTA1 corepressor complex, and its overexpression repressed ER $\alpha$  transactivation functions in a HDAC-dependent manner, proposed to result in the acquisition of the ER $\alpha$ -negative phenotype and increased aggressiveness in breast cancer cells.

A short form of MTA1 protein was subsequently identified and named MTA1s (Fig. 2a) [16]. MTA1s is a splice-variant of MTA1 and contains an ER-binding motif (nuclear binding motif) without any nuclear localization signals at the C-terminus. This protein localizes in the cytoplasm where it sequesters ER $\alpha$ , resulting in the prevention of ligand-induced nuclear translocation of ER $\alpha$  and of stimulation of the malignant phenotype of breast cancer cells. This suggests that the regulation of the cellular localization of ER $\alpha$  by MTA1s may represent a mechanism for redirecting nuclear receptor signaling by nuclear exclusion. MTA1s has also been shown to associate with casein kinase I-gamma2, which is an estrogen-responsive kinase [58].

MTA3 is the latest addition to the MTA family. It was identified as an estrogen-dependent component of the



**Fig. 2** Roles of MTA family in the carcinogenesis and cancer progression. Schematic presentation of the main functions of MTA family proteins. **a** MTA1 protein is included in NuRD complex that represses the transactivation function of estrogen receptor (*ER*)  $\alpha$ , rendering breast cancer cells more phenotypically aggressive. MTA1 protein in NuRD complexes is one of the first downstream targets of c-MYC function, and it is essential for the transformation potential of c-MYC. MTA1s is a splice-variant of MTA1 that localizes in the

cytoplasm where it sequesters *ER* $\alpha$ , resulting in the prevention of the ligand-induced nuclear translocation of *ER* $\alpha$  and stimulation of the development of the malignant phenotype of breast cancer cells. **b** MTA3 protein induced by estrogen represses the expression of the transcriptional repressor Snail, a master regulator of “epithelial to mesenchymal transitions”, resulting in the expression of the cell adhesion molecule E-cadherin and maintenance of a differentiated, normal epithelial status in breast cells

Mi-2/NuRD transcriptional corepressor in breast epithelial cells [15]. The absence of MTA3 as well as the absence of ER results in an aberrantly increased expression of the transcriptional repressor Snail, a master regulator of epithelial-to-mesenchymal transitions (EMT). This increased expression of Snail results in reduction of the cell adhesion molecule E-cadherin expression and subsequently changes in epithelial architecture and invasive growth (Fig. 2b). MTA3 is a transcriptional target of ER $\alpha$ , and in the presence of estrogen, ER $\alpha$  directly binds to the MTA3 promoter at the SP1 site in close proximity of the ERE half-site, resulting in stimulation of MTA3 transcription [59, 60]. Thus, MTA3 functions to maintain a differentiated, normal epithelial status in breast cells, which is in stark contrast to MTA1 or MTA1s. Any potential up-regulation of MTA1 may repress MTA3 expression through repression of the ER $\alpha$  function, leading to up-regulation of Snail, down-regulation of E-cadherin, promotion of EMT and consequently an increase in metastatic potential in breast cancer cells. In fact, Mishra et al. [59] reported that MTA3 gene expression was regulated by the endogenous MTA1 and the knockdown of MTA1 resulted in a significant increase in both basal and estrogen-induced promoter activity of the MTA3 gene. Furthermore, Fujita et al. [60] revealed that a transient forced expression of MTA1 lead to loss of MTA3 protein in breast cancer cell lines. Interestingly, the same phenomenon was also observed in ovarian cancer cell line, in which MTA1 overexpression resulted in down-regulation of E-cadherin and MTA3 expression and enhanced expression of the Snail and Slug [34].

The expression of MTA3 inhibits ductal branching in virgin and pregnant mammary glands in MTA3-transgenic mice [61]. This property is in contrast to MTA1-transgenic

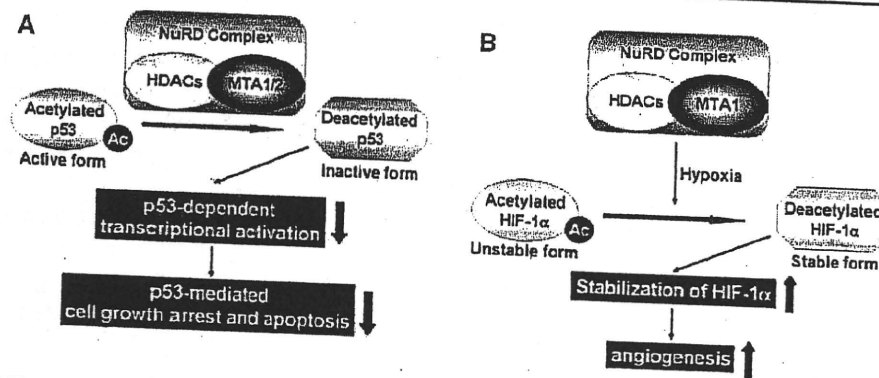
mice, where the inappropriate development of mammary glands results in the development of hyperplastic nodules and mammary tumors, including adenocarcinomas and lymphomas [8, 40]. MTA3 also represses Wnt4 transcription and Wnt4 secretion, inhibiting Wnt-target genes in mammary epithelial cells. This repression of Wnt4 transcription was found to be mediated through a MTA3-NuRD complex, which interacts with the Wnt4-containing chromatin in an HDAC-dependent manner [61].

Although the fundamental functions of MTA proteins are exerted via transcriptional repression by histone deacetylation, a transcriptional activating function has also been demonstrated. Gururaj et al. [62, 63] showed that Breast Cancer Amplified Sequence (BCAS) 3, a gene amplified and overexpressed in breast cancers, was a chromatin target of MTA1, and the transcription of BCAS3 was stimulated by MTA1. This suggested that MTA1 has a transcriptional coactivator function in addition to a corepressor function. A similar finding has also been suggested for mouse *Mta2* protein [64].

#### Deacetylation of non-histone proteins by the MTA family

The protein targets for deacetylation by HDAC via NuRD complexes containing MTA proteins are not only the chromatin histones but also other non-histone proteins. The tumor suppressor gene p53 product was the first non-histone protein that was reported to be deacetylated by MTA protein-containing NuRD complexes. Luo et al. [65] found that the deacetylation of p53 was mediated by a HDAC1 complex containing MTA2 protein. This MTA2-associated NuRD complex interacted with p53 in vitro and





**Fig. 3** Roles of MTA family in the carcinogenesis and cancer progression. Deacetylation of non-histone proteins by MTA family proteins. **a** Tumor suppressor p53 protein is deacetylated and inactivated by both MTA1 and MTA2 proteins in NuRD complexes,

resulting in inhibition of growth arrest and apoptosis. **b** A hypoxia-inducible factor-1 $\alpha$  (HIF-1 $\alpha$ ) is also deacetylated and stabilized by MTA1 protein, leading to angiogenesis

in vivo and reduced significantly the steady-state levels of acetylated p53. Deacetylation of p53 results in an increase of its own degradation through MDM2 and a reduction in p53-dependent transcriptional activation. This eventually leads to the repression of the normal p53 function that mediates cell growth arrest and apoptosis. The same phenomenon was observed between p53 and MTA1. HDAC1/MTA1 complexes possessed deacetylation activity against p53 protein in human non-small cell carcinoma and human hepatoma cells, and the complexes were found to inhibit p53-induced apoptosis by attenuating the transactivation function of p53 [66] (Fig. 3a, b).

Another important non-histone protein that is deacetylated by HDAC1/MTA1 complexes is hypoxia-inducible factor (HIF)-1 $\alpha$ , a key regulator of angiogenic factors [67] (Fig. 3b). The expression of MTA1 is strongly induced under hypoxic conditions in breast cancer cell lines, and MTA1 overexpression enhanced the transcriptional activity and stability of HIF-1 $\alpha$  protein. MTA1 physically binds to HIF-1 $\alpha$  and deacetylates it by increasing the expression of HDAC1, leading to the stabilization of HIF-1 $\alpha$ . These results indicated evidence for positive cross-talk between MTA1 and HIF-1 $\alpha$ , which is mediated by HDAC1 recruitment. They also indicated the existence of a close connection between MTA1-associated metastasis and HIF-1 $\alpha$ -induced tumor angiogenesis. Furthermore, Moon et al. [68] showed that MTA1 increased the transcriptional activity of HIF-1 $\alpha$  and the expression of vascular endothelial growth factor (VEGF), a target molecule of HIF-1 $\alpha$ . Conditioned medium collected from MTA1 transfectants increased angiogenesis in vitro and in vivo. This functional link between HIF-1 $\alpha$  and MTA1 has been demonstrated in clinical samples of pancreatic carcinoma. Using immunohistochemistry and surgically resected pancreatic carcinomas, Miyake et al. [28] examined the expression of HIF-1 $\alpha$ , HDAC1 and MTA1 proteins and suggested that

HIF-1 $\alpha$  expression, which is associated with a poor prognosis in patients with pancreatic cancers, might be regulated by HDAC1/MTA1 complexes. The contribution of MTA1 protein to tumor angiogenesis was also demonstrated in human breast cancers. Using immunohistochemistry, Jang et al. [19] examined MTA1 protein expression and intra-tumoral microvessel density (MVD) in clinical samples of breast cancer and showed that MTA1 expression was significantly correlated with higher tumor grade and higher tumor MVD. The relationship between MTA1 expression and MVD was also observed in HBV-associated HCC [30]. Recently, Yoo et al. [69] experimentally demonstrated that HBV-X (HBx) protein strongly induced the expressions of MTA1 and HDAC1, resulting in those physical link to HIF-1 $\alpha$ . This suggests that positive crosstalk between HBx and MTA1/HDAC1 complex occurs and may be important in stabilizing HIF-1 $\alpha$ , which could play a critical role in angiogenesis and metastasis of HBV-associated HCC [69].

The protein members of NuRD complexes, including MTA1 and MTA2 proteins are co-immunoprecipitated with the ataxia teleangiectasia mutated (ATM)- and Rad3-related protein (ATR) [70]. ATR is a phosphatidylinositol-kinase-related kinase that has been implicated in the response of human cells to multiple forms of DNA damage and may play a role in the DNA replication checkpoint. This fact suggests that MTA proteins may contribute to the regulation of DNA checkpoints.

#### Other possible functions of MTA proteins in cancer

There are other reports suggesting the possible roles of MTA proteins in carcinogenesis and cancer progression. Among them, the most important may be the relationship of MTA1 protein with c-MYC oncoprotein (Fig. 2a). By expression profiling, Zhang et al. [71] identified MTA1 protein as a

target of the c-MYC protein in primary human cancer cells and showed that c-MYC binds to the genomic MTA1 locus and recruits transcriptional coactivators. They also presented data suggesting that MTA1 protein in NuRD complexes was one of the first downstream targets of c-MYC function, essential for the transformation potential of c-MYC, because reduction of MTA1 expression by a short hairpin RNA blocked the ability of c-MYC to transform mammalian cells [71]. There are little data at present concerning the relationship between MTA1 and other important oncogene products such as c-JUN and c-FOS.

As mentioned above, Kumar's group established transgenic mice that overexpressed MTA1 protein and found that the MTA1-transgenic mice showed inappropriate development of mammary glands. These mice also developed hyperplastic nodules and mammary tumors [40]. In this study, the underlying molecular mechanisms were also examined, and the results suggested that MTA1 dysregulation in mammary epithelium and cancer cells triggered down-regulation of the progesterone receptor-B isoform and up-regulation of the progesterone receptor-A isoform, resulting in the up-regulation of the progesterone receptor-A target genes Bcl-XL and cyclin D1 in mammary glands of virgin mice. It would be extremely intriguing and important to examine the HIF-1 $\alpha$ /VEGF expressions and angiogenesis in various organs of the MTA1-transgenic mice, although there are no data concerning these questions at present.

Recently, Molli et al. [72] reported that MTA1/NuRD complexes negatively regulated BRCA1 transcription by physically associating with ERE of the BRCA1 promoter in an ER $\alpha$ -dependent manner and that this repressive effect of MTA1 on BRCA1 expression resulted in an abnormal centrosome number and chromosomal instability. The relationship of MTA proteins with tumor suppressor genes other than p53 and BRCA1 remains to be determined.

Our group showed by the yeast two-hybrid system that mouse Mta1 protein physically linked to endophilin 3 and that the binding of those proteins was made between the SH 3-binding domain of Mta1 protein and the SH-3 domain of endophilin 3 [73]. This suggested that MTA1 protein might be involved in the regulation of endocytosis mediated by endophilin 3.

#### MTA proteins as new molecular targets: clinical implications

On the basis of the available data discussed briefly in this review, it is very likely that MTA proteins have important and critical roles in the genesis and progression of a wide variety of cancers [74]. MTA1 protein can be thought of as a master co-regulatory molecule, strongly and clearly

suggesting the possibility that MTA1 protein (or its gene) could be an excellent molecular target for cancer therapy as well as its use in cancer diagnosis/prognosis. Although studies are not yet available which show the "clinical" efficacy of targeting MTA proteins, several experiments have shown that MTA1 protein (or its gene) could be a molecular target for cancer therapy.

The first studies that suggested the possibility of targeting MTA1 were reported by Nawa et al. [3] and Nicolson et al. [74]. They used antisense phosphorothioate oligonucleotides against *MTA1* mRNA and found a growth inhibitory effect on human metastatic breast cancer cell lines. Since these reports, others have shown that inhibition of MTA1 expression can result in the inhibition of the malignant phenotypes of various cancers, as mentioned below.

Various techniques have been used to regulate MTA1 expression. Using RNAi, Qian et al. [43] inhibited MTA1 expression in a human esophageal squamous cell carcinoma cell line and demonstrated significant inhibition of *in vitro* invasion and migration properties of the cancer cells. The same group further examined the therapeutic value of MTA1 levels in malignant melanoma cells and demonstrated that down-regulation of MTA1 by RNAi successfully suppressed the growth *in vitro* and experimental metastasis of mouse B16F10 melanoma cells *in vivo*, suggesting a promising use of the *MTA1* gene as a target for cancer gene therapy [75].

MTA1s may also be a useful target in the treatment of breast cancer. MTA1s functions as a repressor of ER $\alpha$  transcriptional activity by binding and sequestering the ER $\alpha$  in the cytoplasm [16]. MTA1s has a unique C-terminal 33-amino acid region containing a nuclear receptor-box motif that mediates the interaction of MTA1s and ER $\alpha$ . Singh et al. [76] showed that the MTA1s peptide containing this motif could effectively repress the ER $\alpha$  transactivation function, estrogen-induced proliferation and anchorage-independent growth of the human breast cancer cell line MCF-7. Using an animal model, they also showed the effect of MTA1s peptide in blocking the tumor progression of MCF-7 overexpressing ER $\alpha$ .

There is a good possibility that MTA1 will be a target of immunotherapy. In a review on a model for immunotherapy using a vector, disabled infectious single cycle-herpes simplex virus (DISC-HSV), Assudani et al. [77] proposed that MTA1 is a promising antigen for tumor rejection, because it is greatly overexpressed in many different tumors and is only expressed at lower levels in normal tissues. Their initial studies demonstrated the presence of immunogenic MHC class I-restricted peptides of MTA1. Furthermore, MTA1 was identified as a SEREX antigen, and hence it is likely to be capable of inducing a T-cell response in cancer patients [78].

### Conclusions and future directions

This review has focused on the clinical and biological significance of the newly emerging gene family named MTA, paying particular attention to its relevance to carcinogenesis and cancer progression, such as invasion and metastasis. The fundamental roles of MTA proteins are thought to be transcriptional corepressors that function through histone deacetylation via NuRD complexes, which contain chromatin remodeling and histone deacetylating molecules. Repression of ER $\alpha$  transactivation function by MTA1 protein through deacetylation of ERE chromatin of the ER-responsive genes has been the most extensively investigated, and the data clearly demonstrate that MTA1 expression results in tumor formation in mammary glands and renders breast cancer cells phenotypically more aggressive. In addition, MTA proteins deacetylate non-histone proteins. For example, the tumor suppressor p53 protein is deacetylated and inactivated by both MTA1 and MTA2 proteins, resulting in inhibition of growth arrest and apoptosis. HIF-1 $\alpha$  is also deacetylated and stabilized by MTA1, leading to angiogenesis. Considering the many reports showing the clinical relevance of the expression of *MTA1* mRNA and its encoded protein in a wide variety of human cancers as well as definitive studies showing the molecular and biochemical mechanisms of MTA protein actions, it is likely that MTA proteins, especially MTA1, represent master co-regulatory molecules involved in the carcinogenesis and progression of various malignant tumors. Ultimately this will lead to clinical applications of MTA proteins as a new class of molecular targets for cancer therapy. For example, inhibition of MTA1 expression or function may enhance the chemosensitivity of cancer cells by restoring tumor suppressor function of p53, or it may inhibit tumor angiogenesis by destabilizing the angiogenesis promoting function of HIF-1 $\alpha$ . Moreover, inhibitors of MTA proteins may cooperate with HDAC inhibitors, which are now expected to be a new class of anticancer agents. MTA1 will also be clinically useful for the prediction of the malignant potentials of various human cancers, such as esophageal, gastric and colorectal cancers. Thus, evaluating the expression levels of MTA proteins in individual cases of various cancers may provide clinicians with important clues to prognosis and anticancer therapy.

It will be important to understand the physiological functions and underlying mechanisms of MTA proteins in normal cells, because MTA proteins are also expressed in normal cells and tissues, although at lower levels than found in cancer cells. Physiological roles of MTA1 reported are the followings: (1) MTA1 is thought to play a crucial role in postnatal testis development and spermatogenesis [79, 80], (2) The expression level of MTA1 decreases in mouse brain

in age-dependent manner, which influences the estrogen-mediated signaling pathway during aging [81], (3) MTA1 protein is a direct stimulator of rhodopsin expression [82], (4) MTA1 stimulates hepatic proliferation in vivo and hepatocyte differentiation in vitro [83]. Furthermore, *Caenorhabditis elegans* has MTA1 homologues, *egl-27* and *egr-1*, which are related to embryonic patterning [11, 84] and NURD complex including *egr-1* antagonizes vulval development of *C. elegans*, which is induced by Ras signal transduction pathway [85]. Thus, understanding the physiological functions of MTA proteins will be absolutely necessary to understand the pathological functions of MTA proteins in human cancers. It will be also important to understand MTA1's roles in tissue maintenance via HIF-1 $\alpha$ /VEGF expressions against hypoxic condition.

In conclusion, MTA proteins, especially MTA1, are undoubtedly excellent candidates for therapy and diagnosis/prognosis of human cancers and should be intensively studied for their possible clinical applications.

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# がん診療連携拠点病院に期待される 5大がんの地域連携クリティカルパス

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## SUMMARY

がん診療連携拠点病院に課された5大がんの地域連携クリティカルパスについては、医療現場に混乱が生じている。本稿では、

- ① 行政は「地域連携クリティカルパス」に何を期待しているか、
- ② 地域連携クリティカルパスに寄せる同床異夢、
- ③ 地域連携クリティカルパスの全体像を整理する、
- ④ 連携パスを動かすために必要な仕組みとは、

上記4点に分けて整理を試みた。がんの地域連携クリティカルパスが成立するには医療体制の試行錯誤と大胆な見直しが必要であろう。厚労省の掛け声が質の向上と安心・安全を確保したがん対策の推進につながることを期待したい。

## I 行政は「地域連携クリティカルパス」に何を期待しているか

がん診療連携拠点病院の指定要件(平成18年2月)<sup>1)</sup>として、診療体制に地域の医療機関への診療支援や病棟連携・病診連携の体制の整備が求められ、「地域連携クリティカルパスの整備が望ましい」と明記された。またがん対策推進基本計画(平成19年6月)<sup>2)</sup>では医療機関の整備において取り組むべき施策として個別目標に「すべての拠点病院において5年以内に5大がん(肺がん、胃がん、肝がん、大腸がん、乳がん)に関する地域連携クリティカルパスを整備することを目標とする」とされた。加えて第5次医療法改正の「良質な医療を提供する体制の確立を図るための医療法等の一部を改正する法律」(平成18年法律第84号(改正法)平成18年6月21日交付)に基づく「疾病また5事業ごとの医療体制についての医政局指導課長通知」(医政指発0720001号平成19年7月20日)<sup>3)</sup>では「地域連携クリティカルパスの整備状況」が医療資源・連携等に関する情報として収集されることが記されている。

国として行政が期待する地域連携クリティカルパスは「地域内で各医療機関が共有する、各患者に対する治療開始から終了までの全体的な治療計画(急性期病院から回復期病院を経て自宅に帰り、かかりつけ医にかかるような診療計画であり、医療連携体制に基づく地域完結型医療を具体的に実現するもの)」とあり<sup>1)</sup>、専門的ながん診療機能、標準的ながん診療機能、在宅療養支援機能をもつ医療機関が相互に診療情報や治療計画を共有するなどして連携可能であること(退院後の緩和ケアを含む)を求めている<sup>3)</sup>。

がん対策推進の観点から医療連携、機能分化の前進につながらなければならない。非がん領域を中心にすでに一部の先進的な地域では地域医療ネットワークを構築しつつ、質の高さを追求した地域連携クリティカルパス(以下、連携パス)は稼働し始めており、先進モデルとして医療連携推進の起爆剤となる可能性を秘めている。

## II 地域連携クリティカルパスに寄せる同床異夢

しかし現在のがん診療連携拠点病院，都道府県がん診療連携協議会をみると具体的な連携パスのイメージが描けていない。その原因は2つある。

第一には従来からクリティカルパスの導入に遅れている大学病院が都道府県がん診療連携拠点病院として多く指定されたことである。大学病院は地域医療を超えて教育研究機関として独立した存在であるが，地域行政の展望と指導力の欠如，大学病院の経営努力に追われた拙速な判断ミスと思わざるを得ない。本来がん医療体制は国立がんセンターを中心とした地方がんセンター群（全国がん（成人病）センター協議会）を軸に再編されるべきであった。まさにごん医療体制再構築の前途多難を体現している。

第二には全国的には現段階では行政の思惑に反して医療機関経営者，現場の医療者それぞれの思惑が必ずしも一致していないことである。行政と

しては「診療規模とレベルに合わせて機能分化，役割分担してほしい」と期待するが，医療機関の経営者としては「がん診療連携拠点病院ががん医療の再編のため機動力を持って地域医療の質を支える」というより，地域連携という美名の下に「囲い込み，美味しいところ取りで業績，収益を上げたい」という目先の利益に惹かれてしまっており，他方現場で働く医療者は「院内パスと同じレベルで自分たちの業務に専念し，または専門性を発揮し医療の安全と質の理想を追求したい」という思いを夢見ながら疲弊している。投資とマンパワーが不足しているなかでそれぞれの思いを追求するには無理がある。これらは連携パスの定義と準備が不十分なまま「地域連携クリティカルパス」という響きのよい言葉を拙速に国の指針として入れてしまったことが根本的な問題であったといえる。

## III 地域連携クリティカルパスの全体像を整理する

クリティカルパスとは標準化された診断治療体系に基づいて疾患管理の全体像を可視化，構造化することであり，連携クリティカルパスを平たくいえば医療機能に応じた役割分担の明示である。すなわちその目的は，

第一には医療の質を保証すること，

第二には医療機関の機能分化，役割分担を進めること，

第三にはそれを広く国民に明示することである。

がん診療連携拠点病院はがん医療体制再構築の拠点であり，まずは「標準的治療ガイドラインなどに示された医療について医療機関の機能，規模に応じた役割分担を明示し，連携を調整する」役割がある。医療連携の推進のために筆者が提案したい連携パスの作成指針案は次の通りである（表1）。

すなわち各拠点病院は①医療機関の役割分担表（図1），②共同診療計画表（図2），③私のカルテ（図3），④地域住民に示す医療連携のポスターの4つを作成する。最初は⑤から始め，次に②，③を個別の疾患テーマごとに作成し，各拠点病院が持つ診療ネットワークで運用する。②は連携パスの本

表1 作成の方針

- |  |
|--|
| ● 診療ガイドラインに沿って作成する                     |
| ● 診断・治療施設の役割を明示                        |
| ● 術後の経過観察を各疾患の病期ごとに作成                  |
| ● 診断，治療，外来，在宅，看取りまで                    |
| ● 拠-病-診-看-在-薬-すべての連携を視野に               |
| ● 地域の全医療機関が使えるもの<br>(特定の連携先に限定されない形式を) |



拠点病院	一般病院	かかりつけ医	訪問看護ステーション	居宅介護支援センター	介護療養施設
<ul style="list-style-type: none"> <li>● 精密診断</li> <li>● 手術</li> <li>● 高度な化学療法</li> <li>● 臨床試験</li> <li>● 放射線治療</li> <li>● 定期画像検査</li> <li>● 特殊検査</li> <li>● 専門的全人的緩和ケア</li> <li>● 集学的治療</li> <li>● セカンドオピニオン</li> <li>● 禁煙外来</li> <li>● 院内がん登録</li> <li>● がん医療の情報提供</li> </ul>	<ul style="list-style-type: none"> <li>● 定期健診</li> <li>● 定期検査</li> <li>● 定期画像検査</li> <li>● 麻酔の導入と継続</li> <li>● レスパイト入院</li> <li>● 緊急入院対応</li> <li>● 終末期入院対応</li> </ul>	<ul style="list-style-type: none"> <li>● 初期診断</li> <li>● 画像スクリーニング</li> <li>● 症状管理</li> <li>● 術後補助化学療法</li> <li>● 往診</li> <li>● 在宅ケア</li> <li>● 緩和療法</li> <li>● 看取り</li> </ul>	<ul style="list-style-type: none"> <li>● 病院、かかりつけ医との連携</li> <li>● 訪問介護</li> <li>● 通所介護</li> <li>● 介護保険対応</li> </ul>	<ul style="list-style-type: none"> <li>● 生活療養支援</li> <li>● 介護保険対応</li> </ul>	<ul style="list-style-type: none"> <li>● 療養、身取りの場の提供</li> </ul>

図1 施設の規模・機能による役割分担(内容は試みに例示したもの)

がん種	拠点病院(進ずる病院を含む)	一般病院(有床診療所を含む)	かかりつけ医	訪問看護ステーション	居宅介護支援センター	介護療養施設 居宅
ステージ1	診断、検査 治療 経過観察、対応、ケア	スクリーニング、精密診断 内視鏡的手術、開腹手術 定期観察0ヶ月~12ヶ月ごと、5年間	スクリーニング 術後症状コントロール	日常の指導・管理		
ステージ2,3	診断、検査 治療 経過観察、対応、ケア	スクリーニング、精密診断 開腹手術、標準的補助化学療法 1ヶ月~3ヶ月ごと、5年間 副作用・合併症の対応	スクリーニング 術後症状コントロール、拠点病院(病院)と連携して標準的補助化学療法との継続	日常の指導・管理 副作用・合併症の対応		
ステージ4	診断、検査 治療 経過観察、対応、ケア	スクリーニング、精密診断 開腹手術、標準的補助化学療法、臨床試験を含む高度な化学療法、放射線治療 1週~1ヶ月ごと、拠点病院・かかりつけ医と連携して副作用・合併症に対応	スクリーニング、精密診断 スクリーニング 術後症状コントロール、拠点病院(病院)と連携して治療の継続	日常の指導・管理、拠点病院(病院)と連携して副作用・合併症に対応	病院、かかりつけ医との連携対応 訪問看護、通所介護 介護保険対応	生活療養支援 介護保険対応
緩和医療	診断、検査 治療 経過観察、対応	必要時 連携して症状緩和治療、麻酔の導入と継続 緊急時入院対応、緩和ケア専門入院対応、終末期入院対応	必要時 連携して治療の継続 入院と連携 在宅ケア			療養の場、看取りの場の提供
ケア				かかりつけ医との連携 通所介護 介護保険対応	生活療養支援 介護保険対応	療養の場、看取りの場の提供

\*〇△のときは拠点病院の誰それに連絡して対応する」というような安全と質保証のポイントを決めておく

図2 役割分担表

共同診療計画表 (医療者用)

図3 共同診療計画表 (医療者用)  
術後パスであれば最低限必要な診察や検査、化学療法パスであれば投与計画(間隔など)、標準的な診療計画を提示する。

5大がんの連携パス 私のカルテ

図4 5大がんの連携パス 私のカルテ  
セルフマネジメント用パスを連携に利用、患者がいつも持参することによりどこの施設でも治療経過がわかる。

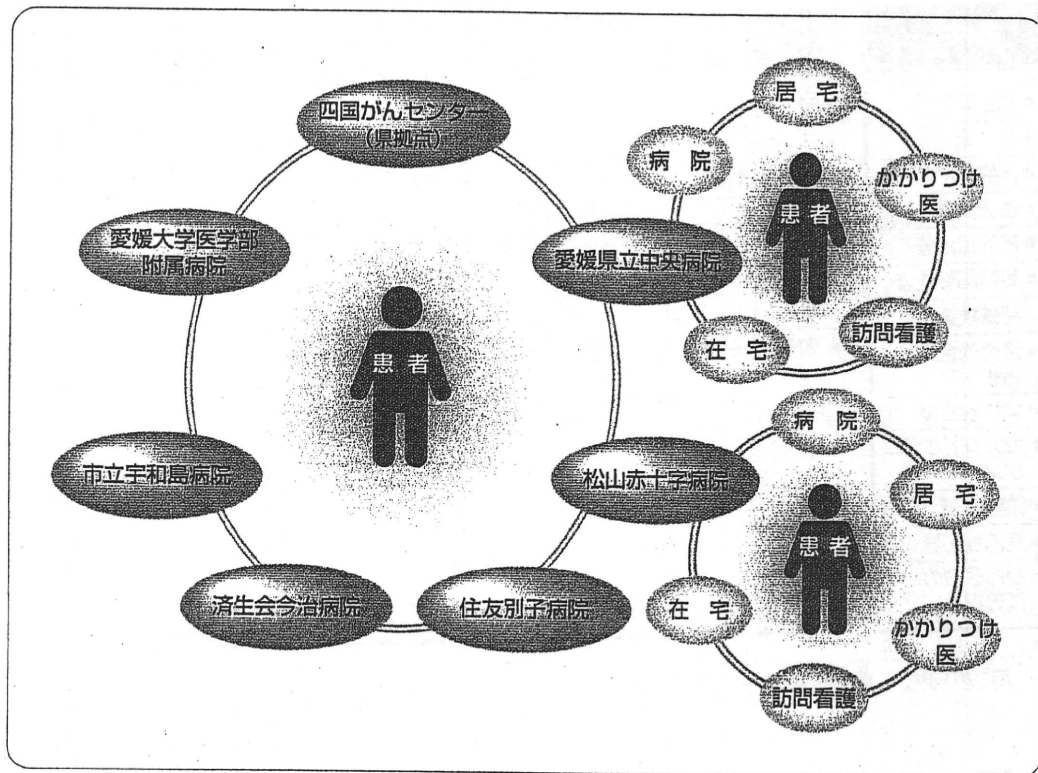


図5 2つのネットワーク構築(愛媛県の場合)  
がん診療連携拠点病院間のネットワーク(左)と各拠点病院が持つ診療ネットワーク(右)。

体に相当し「術後フォローアップ診療計画表」,「化学療法スケジュール表」などがあげられる。③は連携の情報共有ツールとして欠かせない。④は医療連携を広く国民に明示するという意味で患者の受療行動を是正する重要なポイントである。さらに都道府県がん診療連携協議会などで整合性を図

りつつ地域の医療整備状況に合わせていく作業が必要である(図5)。仮にこの指針に則ればまず国全体としてプロトタイプとなる医療規模,機能別役割分担表の提示が求められるであろう。筆者としては今後早い時期にそのプロトタイプを提示したいと考えている。

## Ⅳ 連携パスを動かすために必要な仕組みとは

### 1 地域連携会議, 連携先医療機関との会議

連携する医療機関同士の交流は重要であり,問題意識の高い医療機関との直接の意見交換から医療連携を開始することが出発点である。連携パスの稼動に成功している大腿骨頸部骨折,脳卒中パスなどの先進事例などにその重要性は示されている。しかし連携パスを特定の医療機関間の囲い込みではなく地域連携として成立させるためには直接の医療機関間交流には左右されない地域医療と

してのシステムアップを視野に入れなければならない。拠点病院としての地域医療への責任が問われるべき課題である。

### 2 医療連携室の拡充

医療連携室は外部医療機関との連携の窓口になるだけでなく,医療連携の調整機能を発揮する必要がある<sup>4~6)</sup>。四国がんセンターの「がん相談支援・情報センター」では医療連携室とがん相談支援・情報部門が統合されており,患者の相談対応,

退院支援から医療連携まで広く対応している<sup>7, 8)</sup>。がん診療連携拠点病院の指定要件に示された「がん相談支援センター」<sup>1)</sup>が地域がん医療の quality management を担う必要がある。地域連携クリティカルパスの実働を保証する存在として「連携コーディネーター」の確立と育成を図る必要がある。ここで提案する「連携コーディネーター」はある場合には患者に対して医療者の通訳となり、ある場合には患者の代弁者となり、医療連携の質と安心・安全を保証する存在である。いわば介護保険におけるケアマネジャーに相当する。今後地域

連携クリティカルパスを稼働させるためにはとくに個々の患者ごとの連携を丁寧にコーディネートすることが重要となってくる。現在の医療連携室の枠組みを越えなければならないであろう。職種としての「連携コーディネーター」は看護職をベースとし、日常的に外部にも出向いて活動する。われわれは愛媛医療圏をフィールドとする地域連携モデル実験として「連携コーディネーター」の可能性とあり方を模索し、今後発言していきたいと考えている。

## おわりに

医療機関におけるマンパワーの不足は決定的であり、今は医療現場に多大の負担を強いている。地域連携クリティカルパスは医療現場の必要から発生したものであり、厚生行政の方針として示された医療提供体制の再構築について方向性は正しい。しかし地域連携クリティカルパスの成立にはすこし時間を要する。現場がその解決を図り、工

夫を凝らすにはまだ院内の体制は不備であり、たとえば病棟中心の看護体制などの規制も障害となっている。医療機関の体制に関する試行錯誤と大胆な見直しが必要であろう。今の医療界は変革の時代である「劇的に破壊しつつ大胆に構築する」ことができるかどうか、医療行政、拠点病院の姿勢が問われている。



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# 肺がんの地域連携とクリティカルパス

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## SUMMARY

肺がん術後地域連携クリティカルパスの成功には、クリティカルパスの作成もことながら、個々の症例の連携を重ねて連携の網を形成しておくこと、医療の可化・向上の視点を見失わないことが重要である。また、医療分担の明確化は患者のならず医療者にとっても福音となり得る。今後、地域連携クリティカルパスが切磋磨かれ、一病院に留まらず、地域で共有化されることが望まれる。

## はじめに

第5次医療法改正、がん対策基本法の制定とがん対策推進基本計画の策定など矢継ぎ早の法整備により、地域医療連携とそのクリティカルパス化がにわかに脚光を浴びている。その目的には、よりわかりやすく良質な医療の提供があるだろうし、限られた医療資源をいかに効率よく再配分して医

療体制を構築するのか、といった、マンパワー不足がちな医療者にとって切実な問題も含まれている。本稿では、肺がん術後地域連携クリティカルパス施行2年半の経験から、肺がんの地域連携構築とクリティカルパス化について述べたい。

## I クリティカルパス化によって得られるもの

クリティカルパス化することによりよくなることは、①医療内容が可視化され、②ほかと比較可能となることから、③内容の改訂を余儀なくされるようになる。そして、文書化・定式化することで、④普段当然と思っていた事柄の曖昧な部分が明確化される。また、⑤多忙な外来でつい忘れてしまいがちな検査などの定期チェックが可能となる。最後に、医療者にとって大切なことだが、⑥医療分担の必要性を患者とシェアできることである。

たとえば、当科では1999年に院内最初のクリティカルパスである肺切除術パスが策定された。その目的は大変実務的で、胸部外科部門で多忙であった筆者をどうにか助けてくれようという、優

しい看護諸姉の研究の賜であったが、初期の目は何であれ、ここで問題は定式化された。在院数にドレナージ期間が正比例していることが看研究によって判明し、手術を改善する必要性(胸時肺瘻閉鎖法の再検討)が結論された。抗生も数種類、長い場合は1週間以上も投与されたが、パス化にあたってCEZ一剤に限り、現在術前1回投与まで減少した。これとともに、手時のドレッシングも見直された。それまで組織一部で独自になされていた医療が、可視化・文化されることによって、患者のみならず医療者からも広く評価を受けることとなった。そして、問の解決に際してはEBMを念頭に置く習慣が