Bone morphogenetic proteins in development and progression of breast cancer and therapeutic potential (Review)

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Abstract. Bone morphogenetic proteins (BMPs) belong to the TGF-ß superfamily, which plays important roles in foetal and postnatal development and also maintains the homeostasis of various tissues and organs. Due to the critical role played by BMPs in bone formation and bone turnover, the implication of these molecules in bone metastasis has been intensively studied over the past decade. BMPs have been implicated in the development and progression of solid tumours, particularly the disease-specific bone metastasis. In breast cancer, a tumour type which most commonly metastasizes to bones, aberrations of both BMP expression and their signalling have been recently demonstrated. These aberrations have certain correlations with the development and progression of the disease. Recent in vitro studies have also demonstrated that BMPs can regulate a range of biological functions of breast cancer cells. Targeting BMPs or BMP signalling may provide novel therapeutic approaches for breast cancer. In the current review, we discuss the present knowledge on BMP abnormalities and their implication in the development and progression of breast cancer, particularly in the disease-specific bone metastasis.

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1. Introduction

Bone morphogenetic proteins (BMPs) are members of the TGF-ß superfamily, which participates in the development and homeostasis of diverse tissues and organs through regulating cellular differentiation, proliferation, apoptosis and motility. BMPs exert their effects through a heteromeric receptor complex, which comprises of two types of serine-threonine kinase transmembrane receptors. The Type-I receptors include Activin receptor-like kinase-1 (ALK-1), BMP receptor type IA (BMPR-IA, also known as ALK-3), BMP receptor type IB (BMPR-IB, or so-called ALK-6), ALK-4, ALK-5 and activin A receptor type I (ActRI). The Type-II receptors include BMP receptor type II (BMPR-II), activin A receptor type IIA (ActRIIA) and activin A receptor type IIB (ActRIIB). Upon binding to BMP ligands, the Type-II receptors then phosphorylate the glycine-serine (GS) domain of Type-I receptors. This leads to the recruitment of the pathway-restricted Smads (R-Smads, Smads1, 5 and 8) to the complex. The intracellular signaling complex of R-Smads is then translocated into the nucleus after binding with Smad-4, which leads to the regulation of BMP responsive genes. This pathway is known as the Smad-dependent pathway, in which Smad-6 and -7 act as inhibitory regulators to the signalling. The other pathway is known as the Smad-independent pathway, in which the mitogen-activated protein kinase (MAPK) pathway and the RAS pathway, may be involved (1).

BMPs are key factors in regulating bone formation and bone turnover and have been recently shown to play a pivotal role in tumour development, progression and bone metastasis (1). Breast cancer is the most common cancer in females in the UK and the USA and is the second leading cause of death from cancer. Its leading metastatic site is bone. The major bone lesion in breast cancer is osteolytic, which leads to bone pain, fractures, spine cord compression and hypercalcaemia. These morbidities severely impact the quality of life of the patients. The attention to BMPs and their role in breast cancer has increased as substantial progress has been made in this area. The present review summarises the current knowledge on the roles played by BMPs in breast cancer and their therapeutic potential.

2. BMPs and tumour biology of breast cancer cells

The roles of BMPs in the biology of breast cancer cells have been intensively investigated over the past decade. It is now...
known that BMPs are extensively involved in the regulation of cellular functions of breast cancer cells, ranging from cell growth and death, cell migration, invasion and epithelial-mesenchymal transition (EMT).

Cell cycle and proliferation. BMPs are able to regulate the growth of breast cancer cells. However, the nature of cell response is influenced by the individual BMP, namely, with some BMPs having an inhibitory effect on proliferation of breast cancer cells, while others show a reverse effect (Fig. 1A). For example, BMP-2 and BMP-6 inhibit the proliferation of breast cancer cells (2,3). On the other hand, some BMPs may indirectly promote the proliferation of breast cancer cells, such as BMP-4 which has a synergistic effect on the proliferation of breast cancer cells induced by fibroblast growth factor (FGF), epidermal growth factor (EGF) and hepatocyte growth factor (HGF) (4). This contrasting effect on proliferation of breast cancer cells was clearly demonstrated in a recently published study, where BMP-7 promoted proliferation of MDA-MB-231 and BT-474 cells, but showed an inhibitory effect on the other breast cancer cell lines tested (5).

BMPs can also co-regulate the growth of breast cancer cells induced by non-cytokine agents. For example, BMP-2 inhibits estradiol-induced proliferation of breast cancer cells, via up-regulation of cyclin kinase inhibitor, p21 which in turn inhibits the estradiol-induced cyclin D1-associated kinase activity (6). The up-regulation of p21 by BMP-2 can also prevent EGF-induced proliferation of breast cancer cells (MDA-MB-231) (7). BMP-2 has a direct anti-proliferative effect on tumour cells at a very high concentration (1 μg/ml) in vitro (8). It is interesting to note that the regulation of p21 expression by BMP-2 was mediated by Type-I receptors, Smad-1 and Smad-4. In MDA-MB-468, which only expresses Smad-1, BMP-2 fails to induce p21 and inhibits cellular proliferation (9). BMP-6 inhibits proliferation and induces cell cycle arrest at G0/G1 stage in oestrogen-insensitive breast cancer cells (MDA-MB-231) (2).

To the same BMP protein, different breast cancer cell lines may have a different response. For example, MDA-MB-231, an ER-negative tumour cell line, responds to recombinant human BMP-2 with a greatly reduced proliferation, in comparison to the ER-positive MCF-7 cells (3). This suggests that the oestrogen receptor (ER) status has a bearing on the cell response to BMPs. This link is further strengthened by a study which revealed that BMP-6 and BMP-7 could inhibit oestrogen-induced mitosis of ER-positive breast cancer cells (10).

Apart from the ER status affecting the function of BMPs in breast cancer cells, the BMP receptors must be critical for this diversity due to their essential role in mediating BMP signalling. Recent studies have highlighted certain BMP receptors mediating contrasting effects in breast cancer cells. Overexpression of a domain-negative BMPR-II in breast cancer cells is able to interfere with the phosphorylation of Smad-1 by BMPR-II, leading to an arrest of the cancer cells at the G1 phase of the cell cycle. This suggests that coupling between BMPs and BMPR-II has a significant role in controlling the proliferation and survival of breast cancer cells (11). A domain-negative Type II TGF-β receptor (dnTβRII) could eliminate the anti-proliferative effect of BMP-2 in breast cancer cells by preventing the phosphorylation of Smad-1 (12). One of the Type I receptors, BMPR-IA (ALK-3) has been recently shown to be involved in the activated Smad pathway which contributes to development and progression of breast cancer at primary and secondary sites (13). While BMPR-IA and BMPR-II play positive roles for BMP induced proliferation and aggressiveness in breast cancer cells, another Type I receptor, BMPR-IB has been indicated as a negative regulator (14).

Taken together, BMPs induce diverse effects in breast cancer cells due to the phenotypic profile of BMPs receptors, intracellular signalling molecules and regulatory factors.

Apoptosis. In addition to the pivotal role in the control of cell proliferation and growth, BMPs also play a profound role in regulating apoptosis of breast cancer cells (Fig. 1A). Most interestingly, BMPs can induce a biphasic effect on apoptosis in breast cancer cells depending on the type of BMPs, cell type and experimental circumstances. For example, under routine culture conditions, BMP-2 showed a pro-apoptotic effect in some breast cancer cells (MCF-7), in which the expression of p53 and p21 was increased (15). Under different experimental conditions, BMPs may play a contrasting role in regulation of apoptosis. For example, without supplement of serum, BMP-2 increases the resistance of breast cancer cells
cells do express tenascin-W-coated filters. However, 4T1 mammary carcinoma aggressiveness of breast cancer cell (MCF-7) growth, dominant effect in promoting the motility and invasiveness topic of investigation (Fig. 1B). BMP-2 has shown a pre-regulation of cancer cell motility and invasiveness remains a disseminate and spread. The possible role BMPs play in invasiveness are essential capacities for cancer cells to Motility and invasiveness of tumour cells. Motility and invasiveness are capacities for cancer cells to E-cadherin-mediated cell-to-cell adhesion and prevents breast cancer metastasis through the down-regulation of ßEF1. Higher level of ßEF1 expression is associated with a more invasive phenotype of breast cancer cells (26). Another example is BMP-7, which is able to increase cytokeratin expression and decrease vimentin in breast cancer cells in vitro and in vivo, leading to an epithelial-like phenotype (27) (Fig. 1B).

**BMPs and tumour related angiogenesis.** Angiogenesis is an important event during the development and progression of both primary and secondary tumours. Current knowledge regarding the role of BMPs and angiogenesis is mainly from studies in prostate cancer and bone formation. It has been demonstrated that BMPs, including BMP-2, -4, -6, -7 and GDF5, are capable of inducing angiogenesis. This may be one of the ways by which these BMPs contribute to the process of bone formation (28,29). Experimental evidence suggests that BMPs promote angiogenesis indirectly through up-regulation of the expression of VEGF in both prostate cancer cells and osteoblasts (30,31). The early stage of bone induction by rhBMP-2 can be blocked by the anti-angiogenic agent, TNP-470 (28). This evidence suggests that the control of angiogenesis by BMPs may be, to some extent, coupled with osteoblastic activity. Presently, the understanding of the role of BMPs in angiogenesis in breast cancer is poor. There is very limited literature in this aspect, one study has reported that BMP-2 promotes tumour-related angiogenesis through stimulating p38 MAPK pathway and ID-1 (32). This is obviously a fertile area for explorers.

3. **Crosstalk between BMPs and estrogen signalling**

The involvement of oestrogen in the development and progression of breast cancer is well established. Prolonged exposure to oestrogen in circumstances such as early menarche, late menopause and nulliparity has been considered as high risk to developing breast carcinoma. The effect of oestrogen is largely mediated through two oestrogen receptors, ER-α (ESR1) and ER-ß (ESR2). ER status is correlated with prognosis of patients with breast cancer and is thus a key indicator for selecting endocrine therapies. A possible relationship between BMP/BMP receptor signalling events and ER is an interesting topic. Some of the recent studies have shown some positive correlations.

First, it seems that oestrogen is able to regulate the expression of BMP and BMP receptors. Oestrogen can repress the expression of some BMP receptors, such as BMPR-IA,
BMPR-IB, ACVR2A, and ACVR2B, but has no effect on the expression of ACVR1 and BMPR-II (10). In line with this observation, the expression of some BMPs and BMP receptors in breast cancer tissues has been shown to correlate with ER status. The expression of BMP-7 has been found to highly correlate with the expression level of both estrogen receptor (ER) and progesterone receptor (33). Using enzyme restriction PCR (MSRE-PCR), as well as bisulfate sequencing (BSG), methylation of the human BMP-6 gene promoter was detected in MDA-MB-231; while in MCF-7 and T47D, the BMP-6 gene promoter remained demethylated. In 33 breast tumour specimens, promoter methylation of BMP-6 was detected by methylation-specific PCR. Hypermethylation of BMP-6 was observed in ER-negative cases [16 of 16 cases (100%)], while obviously lower methylation frequency were observed in ER-positive cases [3 of 17 cases (18%)], indicating that BMP-6 expression of ACVR1 and BMPR-II (10). In line with this results. BMP-6 mRNA was detectable in breast carcinoma tissues, its level was reduced in breast carcinoma (18/44) compared with tumour-free resection margins. However, higher levels of BMP-6 mRNA were also found in 8 of the 44 patient samples in comparison with non-tumour margins (43). BMP-2 transcript was decreased in breast primary tumour, in contrast to that of activin-ßA and osteopontin (OPN), which were elevated in these primary tumours (44). Decreased BMP-7 expression in primary tumours was associated with bone metastasis (27). Decreased expression of GDF-9a and BMP-12 was seen in breast cancer compared to matched background tissue and lower expression levels were associated with poor clinical outcome (21,45). A more recent study from the host lab examined the expression of BMP-2 to BMP-7 in a breast cancer cohort using immunohistochemical staining and quantitative PCR. The results showed that BMP-2 and BMP-7 were decreased in breast cancer and correlated with poor prognosis (46).

In contrast to these findings, elevated expression of BMPs has been demonstrated in other studies. For example, BMP-7 was only seen in nuclei of breast cancer cells, but was not detectable in normal breast tissue using immunohistochemical staining (47). BMP-2 and BMP-4 were highly expressed in invasive breast cancer compared to the adjacent normal mammary tissue (16). BMP-4 and BMP-7 were elevated in breast cancer compared to normal breast tissue (48,49). Upregulated BMP-7 expression in primary tumours may correlate with the disease-specific bone metastases of breast cancer (50). BMP-5 was highly expressed in breast cancer and the increasing expression correlated with poor prognosis (46).

Meanwhile, investigations into the expression patterns of BMP receptors and intracellular signalling molecules have also been conducted, but to a rather limited extent. Elevated expression of BMPR-1B was associated with high tumour grade, high tumour proliferation, cytogenetic instability and a poor prognosis in oestrogen receptor-positive carcinomas (51). This suggests that the expression of this type I receptor may associate with the ER status and is regulated by estrogen. The results from the host lab showed a decreased level of BMPR-1B in breast cancer, which was associated with poor prognosis (52).

Activation of the Smad pathway of BMPs (Smad1/5/8) and TGF-ß (Smad2) was revealed in nuclei of breast cancer cells in both primary tumours and bone metastases and similar involvements were also seen in an in vivo model. TGF-ß3 and BMP-2 could promote motility and invasiveness of breast cancer cells (MDA-231-D) in vitro. Moreover, expression of
domain-negative receptors for TGF-β and/or BMPs in the MDA-231-D cells inhibited invasiveness in vitro and bone metastasis in the xenograft model. These results suggest that BMPs as well as TGF-β promote invasion and bone metastasis of breast cancer (13).

In contrast to investigation into clinical breast cancer, studies on the expression of BMPs in breast cancer cell lines appear to be more consistent. Semi-quantitative PCR indicated that BMP-2 and BMP-3 but not BMP-4 and BMP-7, were present in MDA-MB-231 and MCF-7 cells. BMPR-1A, BMPR-1B and BMPR-2 were all detectable in both breast cancer cells at mRNA level. Both transcripts and protein of BMP-2 in MDA-MB-231 cells were decreased in exposure to radiation, whereas the level of BMP-2 in MCF-7 cells was not changed after radiation (3). This provided information for in vitro studies of BMPs in breast cancer.

Although the expression of specific BMPs, such as BMP-2, -4, -6 and -7 in breast cancer is still controversial, the aberrant expression of these BMPs and BMP receptors has been indicated in the development and progression of breast cancer (Table I). The aberrations in the BMP phenotype and signalling in breast cancer, may due to the ER status and self-adjustment by tumour cells themselves according to the needs for development and progression at different stages.

5. BMPs and bone metastasis from breast cancer

Both clinical and experimental studies suggest profound roles for BMPs in bone metastasis of breast cancer. Decreased expression of BMP-7 has been indicated in primary tumours in association with bone metastases. BMP-7 is able to inhibit the growth of breast cancer tumours at primary sites and in bone in vivo (27). Orthotopic implant of tumours with silk scaffolds which were coupled with bone morphogenetic protein-2 (BMP-2) and seeded with bone marrow stromal cells (BMSC), contributed to metastatic spread of breast cancer cells (53). These studies suggest that BMPs are involved in the bone metastasis of breast cancer. On the other hand, lack of BMP antagonists in breast cancer may contribute to the osteoblastic lesions of breast cancer. Conditioned medium (CM) from breast cancer cells (HT-39) could result in an up-regulation of bone sialoprotein mRNA expression in osteoprogenitor cells (MC3T3-E1 cells) and a promotion of their osteoblastic behaviour. This effect could be blocked by addition of Noggin, a BMP antagonist (54). A more recent study also demonstrated that lack of the noggin expression in both breast and prostate cancer cells was associated with osteoblastic activities in the bone metastases. Forced expression of Noggin in an osteo-inductive prostate cancer cell line (C4-2B) reduced in vivo osteoblastic responses induced by its intravenous xenografts, but had little or no influence on bone resorption and tumour growth (55). Unlike Noggin, another BMP antagonist, Gremlin, has been demonstrated to be overexpressed in some human cancers, including breast cancer (56). However, the roles that Gremlin and other BMP antagonists play in coordinating the osteoblastic and osteolytic activities in bone metastatic lesions are far from clear.

CM from breast cancer cells (MCF-7) or prostate cancer cells (LNCaP) could up-regulate osteopontin (OPN) through the protein kinase C (PKC) pathway and mitogen-activated protein kinase (MAPK) pathway. This resulted in inhibition of proliferation and differentiation in osteoblastic cells (57). Another study showed CM from breast cancer (MCF-7) cell-induced apoptosis in preosteoblastic cells (FHSO-6) (58).

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<th>Expression in breast cancer</th>
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<th>Effect on in vivo tumour</th>
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<tr>
<td>Primary tumour</td>
<td>Bone metastasis</td>
<td>Proliferation</td>
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BMP and BMP signalling in breast cancer. ↓, stands for decrease or inhibit; ↑, stands for increase or promote; –, stands for no effect; blank represents unknown.

Table I. BMP and BMP signalling in breast cancer.
6. Perspectives and therapeutic potential

Aberrant expression of BMPs and BMP signalling has been implicated in breast cancer and disease-specific bone metastasis. The expression of some BMPs in primary tumours may have predicting potential, such as that expression of BMP-7 may be associated with bone metastasis and decreasing levels of Noggin may indicate bone metastasis and more osteoblastic activities in bone lesions.

The more recent studies have demonstrated activation of BMP signalling in both breast primary tumours and bone metastases, which contribute to aggressiveness of tumour cells, and development of bone lesions. Lack of Noggin in both breast and prostate cancer cells correlates with their active osteoblastic feature. In the in vivo bone tumour model, Noggin, an antagonist of BMPs, has been shown to prevent bone metastasis by inhibiting both osteoblastic and osteolytic processes. These findings collectively indicate a promising therapeutic value for BMPs and their antagonists in the management of bone metastases. BMPs and their signalling play a profound role in the development, progression and metastasis of breast cancer. Further investigation will elucidate the mechanisms underlying the involvement of BMPs in breast cancer. It will expand current understanding for the pathogenesis of breast cancer and may provide clues for developing novel therapies in managing advanced diseases.

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References


