Application of a three-dimensional (3D) breast cancer model to study macrophage polarization

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Abstract. Knowledge of the tumor microenvironment is crucial for developing an effective strategy to treat cancer. Recently, anticancer therapies targeting macrophages have been intensively investigated. Increased understanding of the importance of the tumor microenvironment has led to the development of three-dimensional (3D) in vitro tumor models. However, established techniques for studying tumor-associated macrophages in vitro are limited. We have previously characterized a 3D breast cancer model consisting of breast cancer cells and fibroblasts cocultured on a silk scaffold. In the present study, the influence of this model on macrophage polarization was investigated. The expression of macrophage markers was studied using reverse transcription-quantitative PCR and flow cytometry. The activity of nitric oxide synthase and arginase in macrophages was also measured. The presented model appeared to induce the polarization of macrophages towards an M2 phenotype. In this 3D tumor model, the in vivo behavior of macrophages could be reproduced. This model may be beneficial for the study of tumor biology and for screening drugs.

Introduction

The tumor microenvironment contributes to cancer growth, invasion and progression (1,2). At all stages of neoplastic development, cancer cells interact with surrounding cellular infiltrates, and vice versa, resulting in the generation of cells with a wide spectrum of activation states. Cancer-associated fibroblasts (CAFs), myofibroblasts, inflammatory immune cells, adipocytes, epithelial cells, endothelial cells and pericytes communicate with cancer cells via cell-cell contact-dependent mechanisms and soluble factors (3-5). Among the immune cells recruited to the tumor development site, macrophages are a particularly relevant population (6).

Macrophages are an indispensable population of immune cells that maintain the homeostasis of the body and are engaged in the course of diseases. They are found in all tissues and are characterized by their complexity and multilateral function (7). Depending on the signals received, macrophages can develop distinct phenotypic programs through a process known as polarization towards either the M1 (classical) or the M2 (alternative) phenotype (7). By contrast to classical macrophage activation, which is associated with the development of an inflammatory reaction, alternative activation is related to inhibition of the effector phase during the immune response (8). Similarly, at the tumor site, tumor-associated macrophages (TAMs) can undergo polarization (9,10). At the early stages of cancer, TAMs assume an M1-like phenotype, but during disease progression, the local signals derived from cancer cells, stromal cells, inflammatory cells and hypoxia promote the polarization of TAMs towards an M2-like phenotype (9). Classically activated M1 macrophages in the tumor environment serve an antiangiogenic and antitumorigenic role (11). At the advanced stage of cancer, M2-like macrophages, which constitute the majority of the TAM population, promote tumor growth by contributing to immunosuppression, angiogenesis and chronic inflammation (11,12).

Current understanding of macrophage polarization is largely based on in vitro studies. Classical activation of macrophages is stimulated by proinflammatory signaling molecules, such as the cytokines interferon (IFN)-γ or tumor necrosis factor (TNF), and can also develop in response to exposure to microbes or microbial products such as lipopolysaccharide (LPS) (7). Such stimulation activates the inflammatory mechanism that leads to increased secretion of proinflammatory cytokines [e.g. TNF, IFN-β], interleukin (IL)-12, IL-6 and IL-1β] and chemokines [e.g., C-C motif chemokine ligand 2 (CCL2), also known as monocyte chemoattractant protein-1 (MCP-1) and C-X-C motif chemokine (CXCL)-10 and CXCL11] and to upregulation of surface molecules, such as major histocompatibility complex (MHC) class II
(I-A/I-E) and CD86 costimulatory molecules (13). Activated M1 macrophages have an enhanced ability to secrete reactive nitrogen species (RNS) and oxygen species, which exert toxic effects on cancer cells (13).

Activation of the macrophage M2 phenotype is stimulated by cytokines IL-4, IL-10, IL-13 and macrophage colony-stimulation factor (14,15). The M2 phenotype is associated with an expression profile of IL-12low, IL-23low and IL-10high and with high production of profibrotic factors such as fibronectin, matrix metalloproteinases (MMPs), IL-1p and transforming growth factor (TGF)-β (16). The most recognized markers of human M2 macrophages are scavenger receptor-1 class A (CD204, also known as SR-AI or MSR1), mannose receptor-1 (CD206), the hemoglobin scavenger receptor (CD163) and macrophage galactose-type C-type lectin (MGL; CD301a) (17-19). In mice, M2 profile markers include arginase 1 (Arg1), resistin-like molecule α (Fizz-1) and chitinase-like protein (19). Additionally, M2 TAMs secrete vascular endothelial growth factor (VEGF), MMPs, epidermal growth factor, TGF-β, IL-10 and CCL2 (20). Immunosuppressive mediators such as IL-10, TGF-β and reactive nitrogen species, which are released by M2 TAMs, suppress T cell proliferation and reduce the antigen-presenting capacity and antitumor response of immunological cells, contributing directly to immune evasion by cancer cells (21).

As macrophages (both M1 and M2) serve a substantial role in tumor development, it is crucial to understand the interaction between macrophages and other cell types in the tumor microenvironment. However, most of the in vitro models generated to investigate the interactions between cancer cells and macrophages are based on two-dimensional (2D) coculture of these cells or the application of conditioned media obtained from cancer cells in 2D culture (22,23). Several studies have examined the regulatory mechanisms underlying macrophage polarization in the tumor microenvironment using a Transwell system, where a porous membrane separates flat cultures of two cell types and cells can thus interact via paracrine signaling (24,25). However, flat 2D cell culture does not mimic the conditions that occur in the body (26). To address this issue, various models of three-dimensional (3D) culture, including microfluidics-based devices, scaffolds and hanging drop technologies, have been proposed (27,28). These models, which enable spatial cell growth, more accurately reproduce the three-dimensional natural microenvironment (27). Due to these approaches, many interactions between cancer cells and cells surrounding the tumor have been demonstrated (29).

Although a number of reports are available (30-32), there is a limited amount of data describing 3D in vitro techniques for studying macrophage polarization in the cancer microenvironment. To address this issue, the present study investigated the influence of a 3D cancer model established in our lab on the status of macrophages.

The 3D in vitro model of breast cancer consists of a spatial coculture of cancer cells and fibroblasts on a silkworm silk scaffold (33). The porosity of the silk scaffold has been optimized to facilitate the growth of cancer cells. Based on the characteristics of the kinetics of cell growth, salt-leached scaffolds with a pore diameter of 250-500 μm were selected to generate a 3D cancer model. To describe this model, the drug cytotoxicity, growth kinetics, morphology and gene expression profile of cocultured cells was investigated and these profiles were compared with cells cultured in a 2D system and in monoluculture on the 3D scaffold. Culture in the 3D coculture system caused a modification of the cells' morphology and significantly increased the production of extracellular matrix (ECM) (33). The interaction of cocultured cells and their spatial growth also induced cellular changes related to epithelial-mesenchymal transition (EMT) and cancer-associated fibroblast markers (33). Moreover, dynamic culture conditions were recently introduced to the 3D model. The implementation of culture medium flow enabled the induction of shear forces into the system which more closely reflects the conditions observed in vivo (34).

In the present study, a functional assessment of the obtained 3D model of breast cancer was performed through analysis of the effect of its microenvironment on macrophages. The conditioned media collected from the 3D cancer model and control 3D cell cultures were used to stimulate macrophages and the phenotypic status of the macrophages was then examined. By using a silk-based 3D breast cancer model the in vivo behavior of macrophages could be reproduced.

### Materials and methods

Establishment of a 3D cancer model and preparation of conditioned media (CM). The EMT6 mouse breast cancer cells and NIH3T3 mouse fibroblasts were obtained from American Type Culture Collection and modified to express green fluorescence protein (GFP) and far-red fluorescence protein (turboFP635), respectively (EMT6/GFP and NIH3T3/635, respectively). The indicated cell modifications were described previously (33). Cells were maintained in DMEM (Sigma-Aldrich; Merk KGaA) supplemented with 10% FBS (Sigma-Aldrich; Merck KGaA) and 80 μg/ml gentamycin (KrKa, d.d. Novo Mesto). Cells were grown at 37°C in a humidified atmosphere containing 5% CO2.

Bombyx mori silkworm cocoons were obtained from the Institute of Natural Fibers and Medicinal Plants (Poznan, Poland) and the silk fibroin solution was extracted as described previously (33). Porous scaffolds were prepared with a salt leaching technique by pouring 0.5 ml silk fibroin solution (~8% w/v) into a polyethylene container 2 cm in diameter and adding 1 g sodium chloride crystals (Thermo Fisher Scientific Inc.) of size 250-500 μm (33). Sodium chloride of the defined particle size was obtained by sieving through 250 and 500 μm test sieves (33).

A total of 3x10^5 EMT6/GFP and NIH3T3/635 cells were seeded on the scaffold at a ratio of 1:9, respectively, as described previously (33). After 2 days, 3x10^5 EMT6/GFP cells or NIH3T3/635 cells were seeded on separate scaffolds. After 7 or 5 days, scaffolds of cocultured and monocultured cells of both lines were transferred into fresh wells of 24-well plates with 3 (unless otherwise specified) scaffolds of each culture type in each well. Next, the cells were incubated at 37°C in fresh medium for 24 h under standard culture conditions. After 24 h, CM was collected and filtered through 0.45 μm filters.

### Macrophage activation.

A total of 1x10^6 J774 cells (American Type Culture Collection) were seeded in 6-well plates and cultured at 37°C for 24 h in DMEM medium supplemented...
with 10% FBS and 80 µg/ml gentamycin. The following day, the medium was removed and replaced with 1.7 ml of CM and incubated at 37˚C for an additional 24 h (unless otherwise specified). J774 cells cultured in 1.7 ml of DMEM medium was the negative control. The sensibility control (positive control for cell reactivity) was J774 cells cultured in culture medium supplemented with mouse IL-4 and IL-6 at a final concentration of 50 ng/ml each.

Fig. 1 provides a schematic representation of the experimental protocol consisting of the establishment of 3D cell cultures, CM preparation and macrophage activation. RNA isolation and reverse transcription-quantitative PCR (RT-qPCR) analysis. Macrophages were stimulated for 12 h with collected CM. Next, RNA was isolated from J774 cells using TRI reagent (Sigma-Aldrich; Merck KGaA) following the manufacturer's protocol. The quality and quantity of RNA were determined spectrophotometrically. High-quality RNA was reverse transcribed using the iScript cDNA synthesis kit (Bio-Rad Laboratories, Inc.) in accordance with the manufacturer's protocol. Expression of Itgax, Mrc1, Ccl24, Retnla, Arg1, Il10, Vegfa, Il6, Cd44, Tgfbl, Tnfa and Ki67 genes was analyzed via RT-qPCR. Reactions were designed to utilize fluorescent hydrolysis probes (Universal Probe Library; Roche Diagnostics) specific towards the expected products. The used gene-specific primers and fluorescent probes are listed in Table I. The primers were designed at the Universal Probe Library Assay Design Center (https://lifescience.roche.com/en_pl/articles/Universal-ProbeLibrary-System-Assay-Design.html) The reaction was carried out on LightCycler 480 (Roche Diagnostics) and using a Probes Master kit (cat. no. 04707494001; Roche Diagnostics) according to manufacturer's protocol. Gene expression was normalized to Tubb (β-tubulin) expression for each sample (Table I). Fold gene expression changes were calculated using the 2^−ΔΔCq method (35). The experiment was repeated at least three times in triplicate.

Quantitation of cytokine/chemokine levels by cytometric bead array. A mouse inflammatory cytometric bead array (CBA) kit (cat. no. 552364; BD Biosciences) was used to determine the concentrations of IL-12p70, TNFα, IFN-γ, MCP-1, IL-10 and IL-6 in CM. Briefly, a mixture of six capture bead populations (50 µl) with distinct fluorescence intensities and precoated with capture antibodies specific for the above-mentioned proteins was added to 50 µl of the sample. Next, PE-conjugated detection antibodies (50 µl) were added to each sample and incubated for 2 h in the dark at room temperature. The unbound antibodies were removed by the addition of 1 ml of wash buffer followed by centrifugation at room temperature (200 x g for 5 min). Next, the captured beads were resuspended in wash buffer (300 µl) and analyzed using a FACSARiaII flow cytometer (BD Biosciences) and FACS Diva v6.1.2 software (BD Biosciences). The amount of each cytokine in the CM was calculated based on the corresponding standard curve with FCAP v3.0 software (BD Biosciences). The experiment was repeated three times.

Analysis of the activated macrophages by flow cytometry. J774 cells were detached from the plate using a cell scraper, washed three times in PBS and analyzed using a FACSARia
flow cytometer (BD Biosciences) with FACSDiva v6.1.2 software (BD Biosciences) or FlowJo v10 software (FlowJo, LLC). The following antibodies were used to analyze J774 cells: PE anti-mouse CD301a (MGL1; Miltenyi Biotec GmbH; cat. no. 130-109-216; 1:10 dilution), PE anti-mouse CD206 (BioLegend, Inc.; cat. no. 141706; 1:200 dilution), APC anti-mouse SR-AI/MSR (R&D Systems, Inc.; cat. no. FAB1797A; 1:1,000 dilution) and PerCP/Cyanine5.5 anti-mouse IA/IE (BioLegend, Inc.; cat. no. 107626; 1:200 dilution). The antibodies were added to the cells and incubated at room temperature for 30 min. PE- and PerCP/Cyanine5.5-conjugated antibodies were excited with a 488 nm laser, and emitted light was collected via 575/26 and 695/40 filters. The APC-conjugated antibody was excited with a 633 nm laser, and emitted light was collected via a 660/20 filter.

Arginase activity. Arginase activity was measured in macrophage lysates by using an Arginase Assay kit (cat. no. Z5030047; BioChain Institute, Inc.) according to the manufacturer’s protocol with minor modifications. Macrophages were activated as aforementioned and following washing with cold PBS and collection of cells with a cell scraper; a 1x10⁶ cells were lysed by adding 100 µl of 10 mM Tris-HCl (pH 7.4) buffer containing 1 mM protease inhibitor cocktail (Merck KGaA) and 0.4% Triton X-100 (Sigma-Aldrich; Merck KGaA). After centrifugation (14,000 x g at 4°C for 10 min), supernatants were placed into a 96-well plate and used for arginase activity assays. Briefly, 10 µl of reaction buffer consisting of the substrate and cofactor was added to 40 µl of the sample to allow the conversion of arginine to ornithine and urea by arginase. Arginase activity was proportional to the produced urea concentration, which was measured by adding 200 µl of the chromogen. The colored complexes were measured using an Elx808™ Absorbance Microplate Reader (BioTek Instruments, Inc.) at a wavelength of 405 nm, and the data were used to calculate the unit arginase activity following the manufacturer’s formula: (ODsample-ODblank/ODstandard-ODwater) x 10.4, where 1 U is the activity needed to convert 1 µMol of L-arginine to ornithine and urea per minute at pH 9.5 and 37°C. The experiment was repeated three times.

Nitric oxide synthase (NOS) activity. NOS activity was quantified in cell lysate using an ultrasensitive colorimetric assay NOS kit according to the manufacturer’s specifications (cat. no. NB78; Oxford Biomedical Research, Inc.). Due to the activity of NOS, nitric oxide (NO) is generated. Upon addition of the kit components, NO undergoes a series of reactions, and the final product of nitrite is measured using Griess reagent. After 24 h of incubation with CM, macrophages were washed with PBS and lysed by the addition of RIPA buffer (Thermo Fisher Scientific, Inc.) supplemented with protease inhibitor cocktail (Merck KGaA). After centrifugation
These genes included those encoding the following proteins: CD11c (Itagx), CD206 scavenger receptor (Mrc1), eotaxin-2 (Ccl24), Fizz-1 (Retnla), arginase-1 (Arg1), and IL-10 (Il10) (Fig. 3). The slight increase in Itagx and Arg1 expression was observed in J774 cells treated with CM collected from the 3T3 monoculture. Medium from the unseeded scaffold did not affect the macrophage phenotype (Fig. 3).

Expression of protumorigenic factors in stimulated macrophages. Stimulated macrophages were analyzed to assess the expression of protumorigenic factors. qPCR showed that the culture of macrophages in CM from the MIX or EMT6 monoculture significantly increased not only Il6 and Tnfa mRNA levels but also Vegfa and Tgfβ mRNA levels in comparison with NC (Fig. 4). The level of Cd44 was significantly increased in macrophages upon stimulation with CM from EMT6 cells in comparison to stimulation with NC. In addition, increased CD44 expression was found after treatment with the MIX sample. The CM collected from the 3T3 monoculture induced an increase in the Tnfa expression level and slightly increased the Cd44 expression level; however, these increases were lower than those induced by MIX and EMT6 CM. CM from none of the tested samples stimulated macrophage proliferation, as indicated by the lack of increased expression of Ki67, a routinely used proliferation marker (36).

Expression of surface markers on stimulated macrophages. Representative results from flow cytometric analysis of macrophages are presented in Fig. 5. Increased expression of Cd301a (MGL-1) was observed on macrophages cultured in the presence of CM collected from the MIX model relative to that from macrophages cultured in CM collected from the other 3D cultures. The lowest expression of Cd301a (MGL-1) was observed on macrophages treated with CM from the 3T3 monoculture. The expression of the CD206 receptor exhibited the smallest difference in activated macrophages. MHC class II (1-A/1-E) expression was lower on macrophages incubated with CM from the MIX model and EMT6 monoculture than on macrophages stimulated with CM collected from the 3T3 monoculture. The expression of SR-AI/MSR was higher on macrophages treated with CM from the MIX model than on cells incubated with CM from the EMT6 and 3T3 monocultures (Fig. 5).

Activity of arginase in stimulated macrophages. Arginase activity was measured in macrophages using a colorimetric

(14,000 x g at 4°C for 10 min), supernatants were collected, and the protein concentration was measured using a BCA protein assay kit (Thermo Fisher Scientific, Inc.) according to the manufacturer's protocol. Next, the assay was performed as described in the manufacturer's protocol with 30 µg of protein. The absorbance was read spectrophotometrically at a wavelength of 540 nm using an ELx808 Absorbance Microplate Reader (BioTek Instruments, Inc.). The experiment was repeated three times.

Statistical analysis. Statistical analyses were performed in GraphPad Prism v5.01 software (GraphPad Software, Inc.) using one-way ANOVA with the Bonferroni test for multiple comparisons. The experiments were repeated at least three times and the data are presented as the mean ± SEM. P<0.05 was considered to indicate a statistically significant difference.

Results

Cytokine secretion in 3D monocultures and the 3D model of breast cancer. Using a CBA assay, the cytokine secretion profiles of CM collected from the 3D monocultures of cancer cells and fibroblasts and from the 3D model of breast cancer were analyzed. In contrast to the MIX and EMT6 monoculture, the 3T3 monoculture produced a significant quantity of MCP-1 (Fig. 2A). The concentration of TNFα in CM collected from the MIX culture was significantly higher than that in the CM collected from the EMT6 and 3T3 monocultures (Fig. 2B). A larger quantity of IL-10 was isolated from the CM of the MIX and EMT6 cultures compared with the 3T3 monoculture. However, these differences were not significant (Fig. 2C). The EMT6 monoculture exhibited a trend towards a larger quantity of secreted IL-6 compared with other cell cultures (Fig. 2D). The secretion of inflammatory cytokines such as IL-12 and IFN-γ was below the detection level in all 3D culture systems (data not shown).

Expression of phenotypic markers in stimulated macrophages. Gene expression in macrophages was analyzed after 12 h of stimulation with CM and compared with that of macrophages cultured with CM collected from unseeded silk scaffolds [negative control scaffold (NCS); Figs. 3 and 4]. Nontreated macrophages were used as an additional negative control (NC). Genes characteristic for the M2 phenotype (9) were upregulated in J774 cells treated with CM from either the MIX culture or the EMT6 monoculture. These genes included

Figure 2. Cytokine concentration in CM. The concentrations of (A) MCP-1, (B) TNF, (C) IL-10 and (D) IL-6 were measured using a BD™ Cytometric Bead Array. The experiment was repeated three times. Data are presented as the mean ± SEM. *P<0.05 as indicated. CM, conditioned media; 3T3, CM from 3D fibroblast monoculture; EMT6, CM from 3D cancer cell monoculture; MIX, CM from 3D culture of both fibroblasts and cancer cells; MCP-1, monocyte chemoattractant protein 1; TNF, tumor necrosis factor; IL, interleukin.
The highest arginase activity among CM-treated macrophages was demonstrated in cells incubated with CM obtained from the MIX model; the next-highest, in macrophages incubated with CM from the EMT6 monoculture and the lowest, in macrophages incubated with CM from the 3T3 monoculture (Fig. 6). However, the differences were not significant.

**Activity of nitric oxide synthase in stimulated macrophages.** The intracellular concentration of nitric oxide was measured as

![Graphs showing expression levels of various genes](image)
a marker of NOS activity. The differences between the macrophages stimulated with CM were not significant compared with the NC macrophages (Fig. 7).

Discussion

Currently, in vitro cell culture is the primary tool used to elucidate cell physiology, intercellular-relationships and the impact of various substances on cellular processes. There is a significant difference between the behavior of cells in vivo and those grown in vitro as monolayers (27). As part of the cell surface adheres to the base of the culture dish, cells do not react appropriately to extracellular stimuli. Therefore, certain signaling pathways are artificially silenced. Unfortunately, these pathways are often crucial for the natural response of cells to changes in environmental conditions in terms of biological processes related to metabolism, growth, differentiation and the ability to spread (for example, focal adhesion) (37). Therefore, various approaches have been proposed to generate conditions that more closely imitate nature. As it enables the spatial growth of cells, 3D culture most closely recapitulates the natural microenvironment (38). Silk as a biocompatible material can be used to generate the platforms for developing 3D models of diseases including cancer (33).

A more valuable in vitro model than cell monoculture is coculture that enables a comprehensive understanding of complex intercellular interactions. Coculture is a type of culture in which two or more cell lines are simultaneously combined in the same culture dish. The combination of various cells naturally present in tissue allows for better modeling of a specific tissue environment. In cocultures, cells can adhere directly to each other and contact is mediated by adhesion molecules or receptors. The cells can also communicate by secreting soluble mediators into the environment. In vitro cocultures are used to understand the mechanisms regulating various complex biological processes and to generate the appropriate models for testing modern drugs (39-41).

Our previous studies were focused on building a tumor model that most accurately reflects in vivo conditions (33,34). The natural microenvironment of tumors, in addition to the cancer cells themselves, includes the connective tissue supporting the stroma, immune cells infiltrating the tumor, extracellular matrix and blood vessels, which all participate in cancer nourishment (42). A tumor is a type of ‘organ’ in which...
To meet these requirements, an in vitro 3D model of breast cancer was developed that consists of breast cancer cells and fibroblasts cocultured on a silk scaffold (33). In this cancer model, both tumor dimensionality and cell heterogeneity are considered. The spatial growth of cells enables interactions among cancer cells, cancer-associated cells and the ECM. These interactions induce alterations in the expression levels of ECM components and markers related to EMT and CAF relative to the corresponding levels in cells cultured in 2D systems and in 3D monoculture systems (33). This model more closely represents the in vivo conditions occurring in the tumor environment than traditional 2D culture and may be beneficial to study tumor biology and to screen drugs (33).

However, as mentioned above, the tumor ‘organ’ is composed of more than cancer cells and fibroblasts and building a more relevant tumor model is necessary to recapitulate the complexity of the system. To control and better understand the model, it is essential to add individual elements gradually. Among the immune cells recruited to the tumor development site, macrophages are a particularly relevant population (43). Thus, further research is required to examine macrophage behavior in the microenvironment generated by the model. It has been reported that cells that can modify macrophages in the tumor microenvironment are tumor-infiltrating lymphocytes (44), tumor-associated neutrophils (45) and CAFs. As in the present 3D model of breast cancer, fibroblasts acquire the characteristics of CAFs, the application of this model was particularly justified. As previous results demonstrated significant differences between cells cultured under 2D and 3D conditions and indicated that 3D culture more closely reflects the effects observed in vivo (33), research has continued only on cells that grew on the scaffolds. Indeed, an increasing number of studies, primarily these that investigate the complex interactions in the tumor microenvironment, are focused on only using a 3D model without researching a flat culture system (30,32).

In the current study, levels of cytokines in the CM obtained from cocultured and monocultured cells was measured. This confirmed that the established 3D model of cocultured breast cancer cells and fibroblasts generated a much more complicated microenvironment compared with that of monocultures of each cell line. Measurement of the concentrations of six cytokines: IFN-γ, TNFα, MCP-1, IL-6, IL-10 and IL-12p70, indicated considerable intersample diversity. For MCP-1 and TNFα, the differences were significant. As fibroblasts are the primary source of MCP-1, the lower concentration of this molecule in CM from the MIX system could be a result of a smaller number of fibroblasts in the coculture. As indicated previously, after 1 week of incubation, cancer cells constituted ~70% and fibroblasts constituted 30% of the cocultured cells (33). However, the finding that the CM of the MIX sample contained the highest level of TNFα did not result from a simple additive effect of each separate monoculture. This effect likely resulted from the mutual interaction of both cell types. The addition of fibroblasts, which express lower levels of IL-10, to the cancer cells with higher IL-10 expression did not decrease the total level of IL-10 expression; by contrast, the MIX sample showed a trend towards higher expression of IL-10. For IL-6, the addition of fibroblasts to the breast cancer cells resulted in a slight decrease in the total level of IL-6 in the MIX sample compared with the EMT6 cell monoculture. Previous results indicated that fibroblasts after coculture exhibited a significantly higher level of IL-6 mRNA than fibroblasts in 3D monoculture (33), suggesting that fibroblasts could be the primary source of IL-6 in the MIX sample. However, measuring the mRNA level did not provide information about the amount of protein. It is worth mentioning that the protein levels of the examined factors in the MIX sample indicated the total amount of proteins expressed by both types of cells during coculture. The mRNA levels of the expressed factors examined in the previous study were measured separately in cancer cells and fibroblasts after their separation from coculture (33). In summary, the data presented in both our previous study (33) and the present study indicate that a 3D tumor model generates a different microenvironment than cancer cells and fibroblasts in 3D monoculture.
The tumor model of the present study was used as a tool to assess tumor biology in terms of macrophage activation. Briefly, CM was collected from 3D monocultures of EMT6, murine fibroblasts (3T3) and a coculture consisting of both cell lines (the 3D breast cancer model, MIX) and used to activate macrophages. Next, the stimulated macrophages were evaluated for possible changes in their activation profiles. To evaluate M1-type activation of macrophages, the level of expression of the major histocompatibility complex type II molecule I-A/I-E was controlled. The activity of NOS was measured alongside the level of TNFα gene expression, which is a proinflammatory cytokine. To monitor M2-type activation, the expression levels of CD206, CD301a and SR-AI/MSR molecules and the enzymatic activity of arginase was measured and the expression levels of the Il6, Mcr1, Ccl24, Retnla, Arg1 and Il10 genes were determined. In these macrophages, the expression levels of the Il6, Cd44, Vegfa, and Tgfb1 genes were also examined, which encode factors essential for the development of cancer (15,21). Moreover, to monitor the sensitivity of the in vitro cultured macrophages to stimulation, a sensibility control was introduced that consisted of medium supplemented with IL-6 and IL-4. The IL-4 cytokine is frequently used to induce M2 macrophage polarization (46). However, M2 macrophages are heterogeneous groups of cells and can be divided into several subtypes (16). IL-6 has been indicated to serve a significant role in the breast cancer microenvironment (47). Thus, to closely mimic the microenvironment of breast cancer and the characteristics of TAMs, macrophages were also costimulated with IL-6 in the present study. This sensibility control allowed monitoring of the reactivity of macrophages upon stimulation; however, it did not constitute a direct positive control in the experimental model, as it was assumed that the collected CM contained a more complex cocktail of secreted molecules than the sensibility control.

Regardless of whether the CM originated from the EMT6 cell monoculture or the MIX, the expression levels of genes responsible for the M2 phenotype and genes encoding factors responsible for the progression of cancer were increased similarly in macrophages compared with those treated with control medium. A particularly large increase in the relative expression of the Arg1 and Vegfa genes was noted. Each of these factors are hallmarks of TAMs/M2 macrophages (9). Moreover, the present results suggested an increase in Cd44 mRNA levels in macrophages stimulated with CM from the EMT6 and MIX cultures in comparison with those treated with a control. CD44 is a transmembrane receptor for hyaluronic acid that plays an essential role in cell adhesion, cell interactions with the ECM and lymphocyte activation (48). It has been revealed that IL-10 and TNFα increases CD44 expression in monocytes (49). Since CD44 serves a crucial role in maintaining monocytes in the circulation during inflammation and mediating their homing to the inflammation site (50), monocytes/macrophages may be recruited into the tumor microenvironment through that same mechanism. However, this hypothesis requires further study.

As the sensitivity of the methods used to assess RNA and protein levels are different, the macrophage stimulation protocol was modified slightly. CM was collected from three scaffolds of each 3D culture type and the time of macrophage stimulation increased to 24 h. The data indicated that the concentration of secreted soluble factors differed between CM from the 3D cancer model and 3D monocultures, which could influence the activation profile of J774 macrophages. Indeed, the highest surface expression of the scavenger receptor SR-AI and the galactose-type C-type lectin (CD301a; MGL1) and the lowest surface expression of the I-A/I-E marker was on macrophages cultured with MIX CM. Moreover, it was observed that CM obtained from the 3D tumor model supported the development of the procancer activity of macrophages. J774 macrophages cultured in MIX CM demonstrated reduced NOS enzymatic activity, which was accompanied by increased arginase activity. Although the indicated hallmarks of the M2 phenotype were not limited to macrophages activated by CM derived from cocultured breast cancer cells and fibroblasts, and the observed differences between differently stimulated groups were not always significant, the MIX sample always induced the most pronounced effects on driving macrophages towards M2 polarization.

As the monocoloncultured cells grew faster than the cocultured cells on silk scaffolds (33), media was collected from the 3D culture systems at different time points. The aim was to compare the influence of CMs collected from a similar total amount of cells from the 3D cancer cell monoculture, fibroblast monoculture and 3D tumor model. Although the tumor model consisted of ~70% cancer cells and 30% fibroblasts at the time of CM collection, it was observed that the stimulation of macrophages by MIX CM was, in some cases, similar to (especially at the RNA level) or stronger than the macrophages' response to CM from the cancer cell monoculture. This finding again indicated that coculture of both cell types generated a novel microenvironment composed of a different cocktail of secreted factors. Under those conditions, it was not possible to determine which of the components was a key factor. It was previously indicated that upon coculture with cancer cells, fibroblasts acquire markers characteristic of CAFs (33). Other studies have demonstrated that fibroblasts (under the influence of cancer cells) secrete factors activating macrophages into TAMs with M2 characteristics (51,52). However, the role of the CAFs in the 3D cancer model regarding macrophage activation requires more advanced study.

The activation profile of macrophages cultured in the presence of conditioned media obtained from the 3D tumor model was changed towards that of a procancer M2 phenotype, in a comparable manner to that suggested in the literature (21,32). Recently, extensive controversy and ambiguity has surrounded the classification and nomenclature of macrophage activation profiles. The precision of the originally proposed distribution of M1 and M2 macrophage polarization, along with experimental guidelines. This new insight into macrophage activation profiles is based on information about the source of macrophages, the definition of activators, and a consistent group of markers describing the activation of macrophages. Authors are striving in this way to develop standards for various experiments. In this context, the presently described model can be a more precise tool allowing for the assessment of processes regarding TAM activation in the tumor microenvironment than models based only on the use of monocultures in 2D or 3D conditions.
The present experiments indicated that the 3D coculture of breast cancer cells and fibroblasts, which mimics the conditions in the tumor microenvironment, constitutes a powerful tool for studying relationships between cancer cells and macrophages. The limitation of the applied experimental process is that the macrophages were stimulated with CMs. In such an experimental set-up, only unidirectional interactions can be induced; only factors secreted by cells cultured in a 3D system could influence macrophages. However, mutual interactions are observed in vivo. The next step will be to assess the relationship between the 3D tumor model and macrophages that will be cocultured in the separate counterparts but with paracrine contact between the cancer model and the macrophages. Moreover, a more advanced experimental setup should enable direct contact between cells in the cancer model and macrophages. As the behavior of macrophages is modulated by numerous factors in vivo, the implementation of the present 3D model of cancer to investigate the interactions between cancer and macrophages in animal studies would be an essential next step. The experience gained in the current work is crucial to the study of a more complex experimental system that allows the examination of interactions between the tumor microenvironment and macrophages. Such a system could also be relevant to the estimation of the effects of stimuli such as chemotherapeutic and immunomodulatory drugs in the tumor microenvironment.

The obtained results are crucial to studying macrophage polarization in a more advanced experimental system that allows the examination of the direct interactions between the tumor microenvironment and macrophages in 3D coculture. This knowledge is essential for developing antitumor therapy based on a macrophage-oriented approach. Moreover, the present study is an example of the application of silk biomaterial to generate useful tools for in vitro research.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

HDK and MK were responsible for study conceptualization and confirmed the authenticity of the raw data. AG, ED and KS performed the experiments. MK, HDK, AG, ED and KS wrote the original draft of the manuscript. AG, MK, ED, KS, AM and HDK analyzed and interpreted the data. HDK, MK and AM reviewed and edited the manuscript. All authors read and approved the final manuscript before publication.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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