Pirfenidone reduces immune-suppressive capacity of cancer-associated fibroblasts through targeting CCL17 and TNF-beta

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Abstract

Various factors in the tumor microenvironment (TME) regulate the expression of PD-L1 in carcinoma cells. The cancer-associated fibroblasts (CAFs) play a crucial role in regulating and rewiring TME to enhance their immune suppressive function and to favor the invasion of the malignant cells. Tumor progression may be retarded by targeting CAFs in the TME. Various studies highlighted the ability of targeting CAF with pirfenidone (PFD), leading to increased efficacy of chemotherapy. However, its potential for the reduction of immune-suppression capacity of CAFs remains to be elusive. Here, we assessed the effect of PFD on the expression of PD-L1 on CAF cells. Besides migration inhibitory effects of PFD on CAFs, the expression level of PD-L1 reduced in CAFs after treatment with PFD. The downstream analysis of released cytokines from CAFs showed that PFD significantly dropped the secretion of CCL17 and TNF-β, where a positive association between PFD-targeted proteins and PD-L1 was observed. These data suggest that the treatment of CAF within TME through the PFD may reduce the acquisition of CAF-mediated invasive and immune-suppressive capacity of breast carcinoma cells.

Key words: cancer-associated fibroblasts; pirfenidone; immune-suppression; PD-L1; microfluidic cancer model

INSIGHT BOX

The cancer-associated fibroblasts (CAFs) play a crucial role in regulating tumor microenvironment to enhance their immune-suppressive function. However, targeting this feature of CAFs remained to show. Herein, we employed an integrated in silico and experimental approach to investigate the therapeutic effects of pirfenidone on the immune-suppressive and cancer-promoting capacity of CAFs. The experiments utilize a 3D microfluidic in vitro model where tumor cells can be co-cultured with CAF cells. Employing this platform and our in silico data analysis, we show how pirfenidone inhibited the acquisition of CAF-mediated invasive and immune-suppressive capacity of breast carcinoma cells via suppression of CAFs activation and secretion of cancer-promoting cytokines. Our findings suggest that pirfenidone targeting CAFs may enhance the efficacy of the immune checkpoint inhibitors in breast carcinoma.
INTRODUCTION

Breast cancer (BC) is the most prevalent cancer in women and remains a significant cause of cancer-associated death, despite attempts to provide effective therapies. Patients diagnosed with TNBC do not benefit from endocrine or targeted therapies, and conventional chemotherapy is still considered the clinical state of the art for this subtype. Immune checkpoint blockade therapy has demonstrated success in various cancers but remains limited in TNBC treatment [1]. Among significant mechanisms responsible for tumour cells chemo/immunoresistance, there is increasing interest in understanding the role of the tumour microenvironment (TME) and its components in response to immune checkpoint inhibition [2–4]. Cancer-associated fibroblasts (CAFs) in the TME play a critical role in the complex process of tumour–stroma interaction. In aggressive BCs such as TNBC, the high abundance of CAFs associated with the aggressiveness of adenocarcinoma [5,6], disease recurrence [6–8], drug resistance [9,10] and lack of immune response [5,11]. Although immunomodulatory role of CAFs, as well as their potential application for therapeutic intervention, especially in the field of cancer immunotherapy, remains to be explored [12]. Thus, given their crucial role of CAFs in tumourigenesis and immunosuppression, in combination with the conventional treatments directed against carcinoma cells themselves, targeting CAFs might be a promising therapeutic approach. The recent approval of pirfenidone (PFD) for the treatment of idiopathic pulmonary fibrosis of lung relies on targeting activated fibroblast and their secretory functions [14].

MATERIAL AND METHODS

Figure 1 illustrates the workflow of this study, initiating from in silico data analysis of cancer genome alteration and gene-drug association of pirfenidone (Fig. 1A), followed by microfluidic-based co-culture of CAF cells and breast carcinoma cells in the device (Fig. 1B) and assessing the influence of pirfenidone on expression and the secretion of PD-L1 and associated cytokines (Fig. 1C).

In silico data analysis

The TCGA-BC genomic information and clinical data were downloaded from cBioportal data portal (https://www.cbioportal.org/) and analyzed under Bioconductor tools in R-Software (version 3.8). A detail description of used packages and related scripts to gene expression and mutation analysis is available at https://bioconductor.org/packages/release/bioc/vignettes/maftools/inst/doc/maftools.html. The protein–protein interaction and drug–protein interaction analyses were performed by using STRING PPI and SNITCH package under Cytoscape software (version 3.7.0).

Generation of CAF cells from normal fibroblasts

The normal fibroblasts (NFs) purchased from Lonza (Cat no: CC-2512) and maintained with FGM-2 Fibroblast Growth Medium-2 BulletKit (Cat no: CC-3132) according to provided protocol. To produce MDA-MB-231 conditioned medium, we cultured MDA-MB-231 (Cat no: HTB-26. ATCC) cells at 100% confluence with serum-free Dulbecco’s Modified Eagle’s Media (DMEM) supplemented with 100 U/ml penicillin and 100 μg/ml streptomycin.
Production of conditioned medium from CAF cells

We produced conditioned media of CAF (CAF-CM) using serum-free DMEM to exclude the effects of growth factors present in serum for the following experiments. The confluent (80%) CAF cells were maintained in serum-free DMEM for 48 hours. In experiments designed to analyze the effects of PFD on CAFs, 100% confluent CAFs were cultured in serum-free DMEM containing 100 μM PFD for 48 hours (CAF-CM + PFD) [18]. The conditioned media was centrifuged for 10 minutes at 1000 rpm after collection and stored at 80°C until use. The resulting conditioned media were used to stimulate BC cells. Proteins secreted into the supernatant were also examined through a cytokine profiling array against 42 targets.

Microfluidic device design and cell culture

The microfluidic tissue culture devices used in this study were purchased from AIM Biotech (Singapore). The devices consist of two media channels located on either side of an extended central region named gel channel (Fig. 1B). The 200-μl collagen of two media channels located on either side of an extended central region named gel channel (Fig. 1B). The 200-μl collagen gel solution (2.5 mg/ml) at pH 7.4 was prepared on ice by the mixture of 20-μl 10x PBS with 4-μl NaOH (0.5 N), 129.2-μl collagen type I (Corning), 10-μl freshly trypsinized and dissociated MCF7 at 6 × 10^5 cells/ml (for 2D studies) or 10-μl cell suspension medium with 50–100 tumor spheroids (for 3D studies), and 22.9-μl deionized water. The collagen gel solution containing the MCF7 aggregates or single cells was then loaded into the central gel region of the device and kept in the cell culture incubator at 37°C and 5% CO2 for 40 minutes to allow gel polymerization via thermal cross-linking. Devices with gel only (no embedded cells, control) were prepared as described above except adding cell suspension. Immediately after gel polymerization, DMEM or CAF-CM media was subsequently introduced to respective media channels. In case of migration assay, the central channel was filled with prepared collagen gel solution, while the side channels were loaded with 2000 cell/120-μl cultured medium.

Culturing single cells and multicellular aggregate in the microfluidic device

In order to obtain cell aggregates with the appropriate size for 3D studies, MCF7 cells were trypsinized and resuspended as individual cells at 100 000 cells/ml in DMEM and cultured for 3 consecutive days onto a 100-mm ultra-low attachment dish (Corning Inc, NY, USA). Aggregates were collected and filtered in two consecutive filtration steps: (i) 40-μm filtration, in order to exclude cell aggregates smaller than 40 μm, and (ii) 100-μm filtration, in order to exclude aggregates larger than 100 μm in diameter, and centrifuged by 250 × g for 5 minutes to separate them from the supernatant. For 2D studies, cells were harvested with Trypsin/EDTA and centrifuged at 200 × g for 5 minutes. Cells were resuspended in growth medium and mixed with collagen type I solution for a final concentration of 6 × 10^5 cells/ml total collagen solution as described above.

Immunofluorescent staining

Cell culture media was removed from the devices, and samples in the microfluidic devices were first rinsed in 1× PBS through the adding 70 μl of PBS into one port and another 50 μl into the opposite connected port of a media channel. Then, the cells fixed with 4% paraformaldehyde (Sigma-Aldrich, St. Louis, MO, USA) for 15 minutes at room temperature. Next, 0.1% Triton-X 100 (Sigma-Aldrich, St. Louis, MO, USA) was added, and the device was incubated for 10 minutes before blocking by BSA 1% (cat no: A5611. Sigma-Aldrich, St. Louis, MO, USA) for 2 hours, followed by staining of cells for α-SMA (1:100, cat no: ab197240. Abcam), Vimentin (1:200, cat no: 677804. Biologend) and PD-L1(1:200, cat no: ab214958. Abcam). The mean fluorescent intensity (MFI) of images was analyzed using Cell-Sense software (Olympus, Japan). The NFs and cancer cells cultured in DMEM are considered as control. To cell morphogenesis analysis, the cell area, circularity and aspect ratio were analyzed using MorphoLibJ in ImageJ software.

Cytokine array

The cytokine profiling of CAF-conditioned media included with and without PFD analyzed against a panel of 42 different inflammatory and cancer-promoting cytokines using the Human Cytokine Antibody Array (cat no: ab133997. Abcam) according to the manufacturer’s instructions.

Statistical analysis

All experiments were performed in three independent technical replicates (n = 3). The results of quantitative experiments were expressed as mean ± SD. Statistical analysis was performed with Student t-test. *P-value ≤0.05 was considered as a statistically significant and ***P-value <0.005 was considered as an extremely significant. Microscopic images are representative images from three independent experiments. The MFI was measured and analyzed using ImageJ software.

RESULTS

Association between CAF derived cytokines with PD-L1

According to the drug-bank information, PFD is an orally active small molecule that may inhibit collagen synthesis, down-regulate production of multiple cytokines, and block fibroblast proliferation and stimulation in response to cytokines. In an in silico analysis, first, we analyzed the drug–protein interaction of PFD (Fig. 2A). As depicted in Fig. 2A, PFD potentially targets IL-10 (score: 0.836), TGFβ1 (score: 0.859), TIPM1 (score: 0.824) and FN1 (score: 0.822) (Fig. 2A). Interestingly, the tissue-specific protein–protein interaction analysis between proteins targets of PFD and immune suppression protein PD-L1 (CD274) showed a positive correlation between IL10, and IL6 with CD274 (PD-L1), IL8 (CXCL8) and TNF-β (LTA) in BC (Fig. 2B). Following these observations, we analyzed alteration frequencies of these proteins across 1085 breast carcinoma samples deposited in TCGA (Fig. 2C). As shown in Fig. 2C, high expression of TNF-β, TGF-β1, IL8, IL6 and IL10 was identified across patients diagnosed with breast invasive ductal carcinoma (IDC) compared with other subtypes such as breast invasive lobular carcinoma and mixed ductal-lobular carcinoma (Fig. 2C). In line with the PPI interaction results, the Pearson correlation analysis shows a positive correlation between IL6, IL8, IL10 and TNF-β with PD-L1 among samples with IDC subtypes (Supplementary Fig. 1). It is well established that tumor stromal cells, particularly CAFs, can reprogram cancer cells in TME through the secretion of various cytokines and induce an immune-suppression TME [19]. In line with these findings, we
analyzed the expression of PD-L1 among samples with low- and high-stromal index (LSI/HSI) according to their ESTIMATE score (Fig. 2D) [20]. As we expected, a significant expression of the PD-L1 (Fig. 2D) was observed among the HSI tumor samples compared with the LSI group, indicating a positive association between the proportion of tumor stromal cells and the expression of PD-L1. Taken together, these data suggest that targeting CAFs and its released cytokines may reduce the immune-suppression capacity of TME through the reduction of PD-L1 expression.

CAFs induce a phenotype switch in cancer cells

To generate CAF cells, we cultured normal human lung fibroblasts with condition medium derived from MDA-MB-231 for 72 hours (Fig. 3A), followed by the characterization of induced cells in terms of expression of the CAF marker alpha-smooth muscle actin (α-SMA) and also the expression of PD-L1. As an expression of α-SMA is one of the hallmarks of CAF cells, we observed a significant expression of α-SMA in cultured fibroblasts with MDA-MB-231 condition medium compared with the cells cultured with DMEM (Fig. 3B and C). Various studies have shown the expression of PD-L1 on CAF cells [5]. In line with these results, we assessed the expression of PD-L1 on generated CAFs at the protein level. As depicted in Fig. 3D, our generated CAFs significantly expressed PD-L1 compared with the NFs.

Numerous in vitro and in vivo studies highlighted the migration and recruitment of CAF cells to the TME [21]. To model this characteristic feature of CAFs, we performed a co-cultured assay in a three channels 3D microfluidic cell culture device (Fig. 4A), where the central channel was filled with collagen type-I, while CAF cells and cancer cells were cultured in one of two side channels (Fig. 4A). Compared with the NFs, the CAFs not only invaded the central channel containing collagen type-I and migrated toward the cancer cell containing channel (Fig. 4B) but also significantly expressed both α-SMA (Fig. 4C) and PD-L1 (Fig. 4D), indicating a trans-differentiation of NFs into CAF.

The phenotypic transformation from an epithelial- to mesenchymal-like is a significant step in invasion and metastasis of carcinoma cells, which can be induced through the tumor stromal cells, particularly CAF cells. To assess the effects of cytokines derived from CAFs on the phenotypic switch of carcinoma cells, we cultured MCF7 cells known as an epithelial-like breast carcinoma with CAF-CM for 72 hours, followed by a shape descriptor analysis. As depicted in Fig. 5, in line with a microscopic observation (Fig. 5A), increase in aspect ratio and reduction of circularity level indicated phenotypic switch of carcinoma cells from epithelial to a mesenchymal-like phenotype acquiring invasive phenotype induced by CAF-CM (Fig. 5B and C. P < 0.001). These results are in line with previous observations showing that CAF cells directly or indirectly able to reprogram cancer cells toward the induction of an aggressive TME.

CAFs-induced expression of PD-L1 on PD-L1 low cancer cells

To determine whether CAFs cells able to induce the expression of PD-L1 on BC cells, we generated tumor aggregate from MCF7 cells, a PD-L1 negative BC cell [22] and cultured with CAF condition medium (CAF-CM) in a microfluidic device for 72 hours (Fig. 6A). The generated aggregates mixed with collagen type I as standard ECM and were loaded in the central channel of the device followed by filling the side channels with standard cell
Pirfenidone reduces immune-suppressive capacity of cancer-associated...
Figure 4. Migration of CAFs. (A) The schematic design of experiments in a microfluidic device by culturing CAFs in one of side channels, while the MDA-MB-231 cells are cultured in opposite channel. The central channel is filled by collagen type-I as a standard ECM. (B) Immunofluorescence image of migrating CAFs toward opposite channel where invasive BC cells are seeded. The CAFs highly expressed both $\alpha$-SMA (green) and PD-L1 (red) in comparison with NF (scale bar: 100 μm). (C and D) Quantitative analysis of MFI of $\alpha$-SMA and PD-L1 expression in CAFs and NF ($n = 3$) ($P < 0.005$).

Besides, the level of other cytokines, including OSM, CSF, IL6, IL8 and CXCL1, was reduced following treatment with PFD. Additionally, the PPI network analysis showed a positive association between PFD targeted cytokines and PD-L1, particularly IL6, CCL17 and CXCL1 (Fig 9I). Taken together, these data suggest that PFD can reduce both invasion and immune suppression capacity of cancer cells in TME by targeting various cytokines in CAF cells.

DISCUSSION

Within the tumor-stromal milieu, CAFs are the most prominent cell type and are known to be critical contributors to tumor progression, metastasis and drug resistance against a broad spectrum of therapies including immunotherapy [2]. CAF directed therapy designed to either eliminate them or potentially reprogram them back to their normal resting phenotype is showing some promise. For example, a recent study on pancreatic adenocarcinoma has demonstrated that targeting vitamin-D receptor by calcipotriol in stromal cells can reprogram pancreatic cell-derived CAFs, resulting in higher intra-tumoural penetration of gemcitabine, tumor shrinkage and improved patient survival [23]. Although numerous studies highlighted various therapeutic agents to target CAFs in different cancers [15, 16, 24, 25], its effects on the reduction of immune suppression potential of CAFs in breast carcinoma remained to show [26]. Herein, for the first time, we show that targeting CAFs with PFD can retard the immunosuppressive capacity of CAFs and, consequently, its PD-L1 expression stimulation on breast carcinoma cells through production blockage of various cytokines including CCL17, CXCL1, TNF-$\beta$, IL6 and IL8 in CAFs. Additionally, through the using microfluidic co-culture platform [27–29], we demonstrated that the treatment of CAF cells with PFD not only reduces the expression of a-SMA in CAF cells but also decreases the migration potential of CAFs and breast carcinoma cells in a time-dependent manner.

A large number of studies highlighted the critical role of these factors in the interaction of the tumor-stromal cell toward the induction of an immune-suppressive TME [5]. Studies on genome and proteome analysis of CAFs showed that CAFs exhibit particular immunomodulatory secretome including but not limited to these factors to reshape TME toward tumor progression and potentially regulate the innate immune suppression in several ways [30–34]. The CCL17 was reported as a key element in the re-education of monocytes and the generation of tumor-associated macrophages that expressed IDO and PD-L1 [35]. Additionally, it has been shown the production and secretion of CCL17 within TME through the CAFs trigger recruitment of Myeloid-derived suppressor cells and T$_{\text{reg}}$ cells to TME promoting an immunosuppressive TME in various cancers [36–39]. Furthermore, Omland and colleagues
Pirfenidone reduces immune-suppressive capacity of cancer-associated

Figure 5. Phenotypic transformation of MCF7 cells cultured with CAF-CM. (A) Microscopic image of MCF7 cells cultured with CAF-CM (right) and conventional culture medium (left), showing a phenotype switch in group of cells cultured in CAF-CM toward an invasive behavior compared with the control group. (B and C) The quantitative results of morphometric analysis of aspect ratio (B) and circularity level (C) in transformed cells (n = 3).

Figure 6. Culturing MCF7-derived MCA with CAF-CM. (A) Immunofluorescent image of MCA cultured with either CAF-CM or conventional culture medium (DMEM) and expression of VIM (red), PD-L1 (green) and nucleus (blue) in these group of cells (scale bar: 100 μm). (B and C) Quantitative analysis of MFI of PD-L1 (B) and VIM (C) in MCAs. CAF-CM significantly induced expression of PD-L1 (n = 3) (P < 0.005).

[40] reported that CXCL12 and CCL17 that secreted from resident CAFs within Cutaneous basal cell carcinoma TME increase tumor progression and immunosuppression.

The immune-modulation role of inflammatory cytokines has been evident extensively.

A large amount of in vitro and in vivo studies documented that the secretion of IL-6 and IL-8 by CAFs positively regulates the expression of PD-1/PD-L1 axis in TME, in which targeting these interleukins may reverse immune resistance [34,41–43]. In this regard, Tsukamoto and colleagues [41] found that combinatorial...
targeting of IL6 and PD-L1 not only improves the infiltration of IFNγ-producing CD4+ T cells in tumor tissues but also enhances the expression of T-cells attracting related chemokines in TME. Consistence with these results, we show that CAF-CM significantly increased immunosuppression capacity of breast carcinoma cells through the expression of PD-L1, in which cytokine profiling and PPI network analysis depicted a positive association between the secretion of IL6, IL8, CCL17 and CXCL1 with PD-L1. In addition to these data and line with previous studies [44], we also show that CAFs able to stimulate phenotype switch in carcinoma cells from a non-invasive to an invasive phenotype in which increased expression of vimentin in non-invasive carcinoma cells by CAFs can trigger EMT program and invasiveness. Additionally, we demonstrated that treatment...
Pirfenidone reduces immune-suppressive capacity of cancer-associated CAFs

Figure 9. Cytokine profiling of CAF-CM. (A) The picture of cytokine array results performed for CAF-CM before and after treatment with PFD. The top left and bottom right dots are negative and positive control of array, respectively. (B–H) The mean pixel density analysis results of differentially released cytokines TNF-β (B) (p < 0.005), OSM (C), CSF (D), CCL17 (E), IL6 (F), IL8 (G) and CXCL1 (H) from CAF treated with (CAF-CM+/PFD) and without (CAF-CM/-PFD) PFD (n = 3). (I). Protein–protein interaction network and correlation analysis between differentially released cytokines and PD-L1.

CAFs cells with PFD decreased the migration potential of both CAFs and breast carcinoma cells modeled in a microfluidic device. Similar observations were reported previously for migratory inhibition effect of PFD on both cancer cells and CAFs in pancreatic cancer [18] and BC [45]. For instance, Ren and colleagues demonstrated that CAF-CM significantly induces the expression of HOTAIR and consequently promotes EMT program in invasive BC in which targeting CAFs with PFD blocks TGF-β1/HOTAIR axis and decreases the migration potential of MDA-MB-231 cells [45].

Taken together, at an in vitro stage, these data suggest that targeting CAFs with PFD may reduce both metastasis and immune suppressive capacity of CAFs in TME. Additionally, the potential application of PFD to deplete the secretion of the various cancer-promoting cytokines and chemokines expressed in different sub-types of BC (Fig 2C) (Fig 9) suggesting the potential of applying PFD in combination with current treatment regimens particularly immunotherapy. Further studies focused on complex models of TME, including genetically engineered mouse model included with immune systems elements, are required to approve the potential application of PFD on improving immunotherapy and chemotherapy efficacy.

SUPPLEMENTARY DATA
Supplementary data is available at IntBIO Journal online.

CONFLICT OF INTEREST STATEMENT
The authors reported no potential conflict of interest.

REFERENCES

FUNDING
This work is supported by the National Health and Medical Research Council of Australia (NHMRC) [grant number APP1143377].


