Carfilzomib modulates tumor microenvironment to potentiate immune checkpoint therapy for cancer

Qian Zhou, Jinxia Liang, Tong Yang, Jin Liu, Bo Li, Yingchang Li, Zhenzhen Fan, Weida Wang, Wensheng Chen, Sujing Yuan, Meng Xu, Qigui Xu, Zhidong Luan, Zhong-Jun Xia, Penghui Zhou, Yadong Huang, and Liang Chen

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(Note: With the exception of the correction of typographical or spelling errors that could be a source of ambiguity, letters and reports are not edited. Depending on transfer agreements, referee reports obtained elsewhere may or may not be included in this compilation. Referee reports are anonymous unless the Referee chooses to sign their reports.)
Thank you again for submitting your work to EMBO Molecular Medicine. We have now heard back from the three referees who evaluated your manuscript. As you will see from the reports below, the referees acknowledge the interest and novelty of the study. However, they also raise a series of concerns about your work, which should be convincingly addressed in a major revision of the present manuscript.

The referees’ recommendations are rather clear, and there is no need to reiterate their comments. Most of their concerns refer to the need to provide further details, clarifications, and controls, and to improve the presentation of the study in order to make the data and the main conclusions easily accessible to the general readers.

We would welcome the submission of a revised version within three months for further consideration. Please note that EMBO Molecular Medicine strongly supports a single round of revision. As acceptance or rejection of the manuscript will depend on another round of review, your responses should be as complete as possible.

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We are aware that many laboratories cannot function at full efficiency during the current COVID-19/SARS-CoV-2 pandemic and have therefore extended our "scooping protection policy" to cover the period required for a full revision to address the experimental issues. Please let me know should you need additional time, and also if you see a paper with related content published elsewhere.

Please read below for important editorial formatting and consult our author's guidelines for proper formatting of your revised article for EMBO Molecular Medicine.

I look forward to receiving your revised manuscript.

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In case you have no data that requires deposition in a public database, please state so in this section. Note that the Data Availability Section is restricted to new primary data that are part of this study.

7) For data quantification: please specify the name of the statistical test used to generate error bars and P values, the number (n) of independent experiments (specify technical or biological replicates) underlying each data point and the test used to calculate p-values in each figure legend. The figure legends should contain a basic description of n, P and the test applied. Graphs must include a description of the bars and the error bars (s.d., s.e.m.).

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9) Our journal encourages inclusion of "data citations in the reference list" to directly cite datasets that were re-used and obtained from public databases. Data citations in the article text are distinct from normal bibliographical citations and should directly link to the database records from which the data can be accessed. In the main text, data citations are formatted as follows: "Data ref: Smith et al, 2001" or "Data ref: NCBI Sequence Read Archive PRJNA342805, 2017". In the Reference list, data citations must be labeled with "[DATASET]". A data reference must provide the database name, accession number/identifiers and a resolvable link to the landing page from which the data can be accessed at the end of the reference. Further instructions are available at .

10) We replaced Supplementary Information with Expanded View (EV) Figures and Tables that are collapsible/expandable online. A maximum of 5 EV Figures can be typeset. EV Figures should be cited as 'Figure EV1, Figure EV2" etc... in the text and their respective legends should be included in the main text after the legends of regular figures.

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- Additional Tables/Datasets should be labeled and referred to as Table EV1, Dataset EV1, etc. Legends have to be provided in a separate tab in case of .xls files. Alternatively, the legend can be supplied as a separate text file (README) and zipped together with the Table/Dataset file.

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13) Author contributions: the contribution of every author must be detailed in a separate section (before the acknowledgments).

14) A Conflict of Interest statement should be provided in the main text.

15) Every published paper now includes a 'Synopsis' to further enhance discoverability. Synopses are displayed on the journal webpage and are freely accessible to all readers. They include a short stand first (maximum of 300 characters, including space) as well as 2-5 one-sentences bullet points that summarizes the paper. Please write the bullet points to summarize the key NEW findings. They should be designed to be complementary to the abstract - i.e. not repeat the same text. We encourage inclusion of key acronyms and quantitative information (maximum of 30 words / bullet point). Please use the passive voice. Please attach these in a separate file or send them by email, we will incorporate them accordingly.

Please also suggest a striking image or visual abstract to illustrate your article as a PNG file 550 px wide x 300-600 px high.

EMBO Molecular Medicine has a "scooping protection" policy, whereby similar findings that are published by others during review or revision are not a criterion for rejection. Should you decide to submit a revised version, I do ask that you get in touch after three months if you have not completed it, to update us on the status.

***** Reviewer's comments *****

Referee #1 (Comments on Novelty/Model System for Author):

The authors have screened FDA-approved drugs and found that Carfilzomib drove IL-4 induced macrophages to express M1 cytokines, enhanced phagocytosis ability and promoted T cell proliferation. In vivo, Carfilzomib reprogrammed TAMs into M1-like macrophages, and promoted CD8+ T cell proliferation or activation to inhibit autochthonous lung cancers in a mouse model. And Carfilzomib synergized with PD-1 antibody to better control autochthonous lung cancers. They further identified that Carfilzomib activated IRE1α to recruit TRAF2, resulting in the activated NF-κB to induce expression of the proinflammatory cytokines. In general, this paper have interesting and important findings, which has proposed that the FDA-approved Carfilzomib can be combined together with anti-PD-1 antibody to improve immune therapy against EGFR mutant-lung cancer. This manuscript can be considered for publication and a revision is encouraged after the authors address the below concerns.

My questions are listed below.

1. Fig. 1: In the mock sample, can Carfilzomib alone affect macrophage survival or expression of these proinflammatory cytokines?
2. Fig. 2: Can Carfilzomib alone affect T cell proliferation driven by anti-CD3/CD28 stimulation?

Can Carfilzomib affect the expression levels of MHC-I and MHC-II and CD80 in macrophages, which are critical for antigen presentation? In Fig.2, CD86 levels were checked, while in the in vivo study shown in Fig. 6, CD80 expression was measured. The authors should include both CD80 and CD86 in Fig. 2 and Fig. 6.

The effector on phagocytosis is enhanced by Carfilzomib treatment from 1% to 3%. This might be due to the limited sensitivity of the assay. Nevertheless, this change is not substantial. The authors are suggested to delete the strong description such as significantly. Similarly, words like "drastic" in the manuscript should be modified.

3. Fig. 4J: Carfilzomib treatment can increase TRAF2 binding to IRE1α. But in this figure, the reduced amount of IRE1α is shown after Carfilzomib treatment. IS this correct?

Fig. 4A The Kir6 effect is shown differently in RAW264.7 (completely) vs BMDM (partially). This should be discussed.

4. Fig. S5C: The in vivo study did not observe difference of CD4+ T cells. However, the ex vivo data in Fig. 2 indeed show that Carfilzomib treatment enhances both CD8+ T cell and CD4+ T cell proliferation. This should be discussed.

5. The major question: The IRE1α-TRAF2 module can explain that Carfilzomib treatment enhances expression of the proinflammatory cytokines in IL-4 induced M2 macrophages. But what is the reason to explain the reduced CD206 or Arg1 expression in these cells?

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The manuscript by Zhou et al presents a novel approach to increase the efficiency of PD-1 ICB in resistant cancer. They utilize a variety of experimental procedures and convincingly prove their points. The conclusions are both new and of high medical
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Minor points:
1) I really advise the authors to have the manuscript corrected by a native speaker or a commercial service for language and grammar.
2) In some parts the manuscript is also quite demanding for readers, who are not specialist to the field, so a bit more explanation on experimental systems in the results section would be helpful for the general readership.
3) Especially the model system of tumor-SN-treated Raw264.7 needs a (short) introduction.

Referee #3 (Comments on Novelty/Model System for Author):

Overall, this manuscript by Zhou et al is interesting and timely. The authors identified a way to reprogram immunosuppressive M2 macrophages toward antitumor M1 cells, which may help to develop a novel immunotherapeutic strategy. The experiments are in general informative and reasonably well designed. However, several concerns on the analysis, presentation and interpretation of data are raised (see below).

Referee #3 (Remarks for Author):

In this manuscript, Zhou et al found that carfilzomib, a proteasome inhibitor (PI), could drive murine and human M2 macrophages to partially exhibit M1-like phenotype and function. By screening FDA-approved drugs, the authors identified three PIs namely carfilzomib, bortezomib and MLN9708 that effectively elicited IL-1β expression in murine bone marrow-derived macrophages (BMDMs). Further examination revealed that carfilzomib-induced M1-like cells (Ci-M1). Important, Ci-M1 exhibited enhanced phagocytotic and antigen-presenting activity, suggesting that carfilzomib treatment at least partially endowed M2/TAM with phenotype and functions similar to M1 macrophages. Investigation into the underlying mechanism of carfilzomib-induced M2 transformation into M1-like cells indicated that this process was dependent on an IRE1α-TRAF2-NF-κB pathway. Therapeutically, systemic carfilzomib treatment could inhibit tumor growth and synergize with PD-1 inhibitors in vivo, an effect that was attenuated in the absence of macrophages.

Overall, reprogramming immunosuppressive/protumor macrophages toward cells with antitumor function is emerging as an attractive notion. The present study is interesting and timely. The experiments are in general informative and reasonably well designed. I have some minor concerns about the analysis, presentation and interpretation of data.

1) Figure 1B: Three red dots are shown but they are not individually and specifically annotated. In addition, what does the x axis stand for? It is also unclear whether carfilzomib, bortezomib and MLN9708 are the only drugs in the library that were capable of activating IL-1β expression in IL4-induced M2 macrophages. Is there other PIs in the library that were not able to induce IL-1β expression? These issues should be clarified.
2) Figure 2A: Although carfilzomib could decrease CD206 and arginase 1 expression, this reduction was marginal. Are the levels of IL-6 and iNOS expression post carfilzomib treatment comparable to that in conventional M1 macrophages? A positive control is recommended for supporting the authors’ claim that carfilzomib could reprogram M2 macrophage into M1-like population.
3) Figure 2C: The gating strategy is unclear. How was the CD86+ and CD206+ population defined? Was it based on isotype-matched antibody staining or fluorescence minus one (FMO) control? This point is also applicable for other flow cytometric plots shown in the manuscript (e.g., Figure S2E and S2L).
4) Figure 2F and 2H: Both assays showed that carfilzomib could enhance macrophage phagocytotic ability, but the phagocytosis efficiencies exhibit a ten-fold difference. This should be clarified.
5) Figure 2L, y axis: Is it CD8+ % or CD4+ %? Figure 3D: Extra asterisks are shown on IL-6.
6) Figure 4: It is very interesting that the kinase activity of IRE1α, instead of the endoribonuclease activity and the activation of XBP1, mediated the carfilzomib induced reprogramming of M2 macrophages. The authors are encouraged to discuss the impact of PIs and ER stress inducers on XBP1-deficient cells.
Referee #1 (Comments on Novelty/Model System for Author):
The comments and advice are listed in the remarks to the authors.

Referee #1 (Remarks for Author):

The authors have screened FDA-approved drugs and found that Carfilzomib drove IL-4 induced macrophages to express M1 cytokines, enhanced phagocytosis ability and promoted T cell proliferation. In vivo, Carfilzomib reprogrammed TAMs into M1-like macrophages, and promoted CD8+ T cell proliferation or activation to inhibit autochthonous lung cancers in a mouse model. And Carfilzomib synergized with PD-1 antibody to better control autochthonous lung cancers. They further identified that Carfilzomib activated IRE1α to recruit TRAF2, resulting in the activated NF-κB to induce expression of the proinflammatory cytokines. In general, this paper have interesting and important findings, which has proposed that the FDA-approved Carfilzomib can be combined together with anti-PD-1 antibody to improve immune therapy against EGFR mutant-lung cancer. This manuscript can be considered for publication and a revision is encouraged after the authors address the below concerns.

Reply: We thank Reviewer for finding our work interesting and important.
My questions are listed below.

1. Fig. 1: In the mock sample, can Carfilzomib alone affect macrophage survival or expression of these proinflammatory cytokines?

   **Reply:** Following Reviewer's suggestion, we have now checked macrophage survival and expression of proinflammatory cytokines (IL-6 and INOS) after Carfilzomib treatment. We found that Carfilzomib slightly inhibited the proliferation of macrophages after being treated for 12 hours, and did not affect macrophage cell viability in 6 hours (Figure 1 for Reviewer). Meanwhile, Carfilzomib alone was able to induce *Il-1β*, *Il-6* and *Inos* expression in macrophage (see new Figure EV2E).

   ![](Image)

   **Figure 1 for Reviewer:** BMDMs or Raw264.7 were treated by Carfilzomib (500 nM) for 6 hours or 12 hours (0 hour means beginning of DMSO administration) and CCK8 assay kit was applied to estimate cell viability.

2. Fig. 2: Can Carfilzomib alone affect T cell proliferation driven by anti-CD3/CD28 stimulation? Can Carfilzomib affect the expression levels of MHC-I and MHC-II and CD80 in macrophages, which are critical for antigen presentation? In Fig.2, CD86 levels were checked, while in the in vivo study shown in Fig. 6, CD80 expression was measured. The authors should include both CD80 and CD86 in Fig. 2 and Fig. 6. The effector on phagocytosis is enhanced by Carfilzomib treatment from 1% to 3%. This might be due to the limited sensitivity of the assay. Nevertheless, this change is not substantial. The authors are suggested to delete the strong description such as significantly. Similarly, words like "drastic" in the manuscript should be modified.
Reply: Thanks so much for these great suggestions. We analyzed T cell proliferation driven by anti-CD3/CD28 stimulation after Carfilzomib treatment. The results indicated that Carfilzomib did not affect CD4 and CD8 T cell proliferation (see new Figure EV2Q & EV2R). We also checked the expression levels of MHC-I, MHC-II and CD80 in macrophages in Ci-M1. We found that Ci-M1 expressed higher levels of MHC-II and CD80 but not MHC-I in comparison to IL-4-activated M2 macrophages (new Figure 2C & 2D & EV2M & EV2N). Besides, both CD80 and CD86 are included in new Fig. 2 and Fig. 5. Following Reviewer’s suggestions, we have now corrected the statements throughout the manuscript. We have now deleted “dramatically” or used “markedly” instead of “dramatically”.

3. Fig. 4J: Carfilzomib treatment can increase TRAF2 binding to IRE1α. But in this figure, the reduced amount of IRE1α is shown after Carfilzomib treatment. Is this correct?

Reply: Thanks so much for bringing this important issue into our attention. We repeated this experiment. The results indicated that expression of IRE1α and the recruitment of TRAF2 by IRE1α were significantly increased and in Ci-M1 (New Figure 4J).

Fig. 4A The Kira6 effect is shown differently in RAW264.7 (completely) vs BMDM (partially). This should be discussed.

Reply: We are sorry for not presenting our result clearly enough. Figure 4A showed that IRE1a knockout severely inhibited expression of IL-1B and IL-6 in BMDM. Figure 4B showed that this inhibition is much milder in RAW264.7 cells. The mechanisms underlying different sensitivity of BMDM and RAW264.7 could be complex. We are not sure of it. But several different mechanisms have been reported to control IL-1B transcription, including transcription factors like Spi-1/PU.1 (Mol Cell Biol 15(1) (1995) 58–68), NF-κB plus C/EBPβ and HIF-1α plus C/EBPβ (J Immunol May 1, 2016, 196 (1 Supplement) 189.14), and epigenetic modifications of its promoter region (Arthritis Rheum. 2009 Nov; 60(11): 3303–3313). The
RAW264.7 cell line was established from a tumor induced by the Abelson murine leukemia virus (Ralph P et al., 1977, J Immunol), which was different from BMDM. Therefore, knockout of IRE1a in RAW264.7 and BMDM, different expression of IL-1B and IL-6, which could be reasonably possible. We have discussed in the page 17, line 432-442.

4. Fig. S5C: The in vivo study did not observe difference of CD4+ T cells. However, the ex vivo data in Fig. 2 indeed show that Carfilzomib treatment enhances both CD8+ T cell and CD4+ T cell proliferation. This should be discussed.

**Reply:** We thank Reviewer for bringing this important issue to our attention. Yes, we saw that peptide-loaded Ci-M1 enhance proliferation of both CD4 and CD8 T cells. However, in Carfilzomib treated lung cancer mouse models, we detected similar amount of CD4 T cells in tumors. We guess that the discrepancy between in vitro and in vivo data could be explained by the complex process of maintaining the homeostasis in vivo. Several factors could affect apparent number of CD4 T cells in tumor: recruitment (CXCL9/10), proliferation (IL-15) and death (IL-2) (J Clin Immunol 2002 Mar;22(2):51-6; Blood 2012 Nov 15;120(20):4246-55.). If the microenvironment of Carfilzomib treated lung cancers release less chemoattractant, fewer CD4 T cells are recruited into the tumor locus. However, the faster proliferation of CD4 T cells could compensate for the lower amount of recruited seed CD4 T cells. Or it could be that CD4 T cells underwent faster apoptosis after proliferation. We thank Reviewer again for pointing out his important issue. We have discussed in the page 19, line 485-496.

5. The major question: The IRE1α-TRAF2 module can explain that Carfilzomib treatment enhances expression of the proinflammatory cytokines in IL-4 induced M2 macrophages. But what is the reason to explain the reduced CD206 or Arg1 expression in these cells?

**Reply:** Thanks so much for this insightful comment. In our opinion, IRE1α-TRAF2 activate NF-κB activity, which is responsible for transcribing proinflammatory
cytokines. This effect could explain that effect of Carfilzomib on expression of proinflammatory cytokines is direct. We guess the effect of Carfilzomib on CD206 and Arg1 is indirect, most probably because Carfilzomib can induce M1 polarization. Indeed, impact of M1 polarization on the expression of M2-related genes have been reported. The negative regulation of M2 genes by M1 polarizing signals are relatively easier to understand. For example, Btk activated by LPS inhibits M2 genes by macrophages (Plos One 9, e85834.); NO produced during M1 polarization inhibits macrophages to transcribe IL-10 gene (Blood 117, 5092-5101). On the contrary, M1 polarization can feedback to upregulate M2 related gene in macrophages. For example: M1 macrophages feature activated glycolysis (Immunity 2015. 42:419–430); Lactic acid, the intermediate metabolite of glycolysis, has been found to promote expression of M2-related genes (Cell Commun Signal. 2018;16(1):54). Therefore, the impact of Carfilzomib on expression of M2-related genes (like CD206 and Arg1) is indirect and more environment-dependent. We have discussed this issue in detail in Page 16-17, line 413-431.

Referee #2 (Comments on Novelty/Model System for Author):

The manuscript by Zhou et al presents a novel approach to increase the efficiency of PD-1 ICB in resistant cancer. They utilize a variety of experimental procedures and convincingly prove their points. The conclusions are both new and of high medical impact.

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through its biological effects, its ability to render TAM inflammatory, the mode of action and the intracellular signaling involved. Then they address the effects of Carfilzomib in vivo in an autochthonous murine lung cancer model and show impressive results, especially in combination with a PD-1 inhibitor. Finally, they show similar effects on human ex vivo differentiated macrophages, which strongly argues for transferability to human application. Hence the combination of proteasome inhibitors my represent a new combination treatment to help all those cancer patients who are unresponsive to PD-1 checkpoint blockade. The only prerequisite would be a sufficient TAM infiltration of their tumor. The authors performed their experiments well examined their assumptions often with different methods. Hence, I consider this a high quality paper from the scientific point of view. Unfortunately, the English language is, in many parts, not sufficient for publication.

Reply: We thank Reviewer for finding our study novel and compelling.

Minor points:
1) I really advise the authors to have the manuscript corrected by a native speaker or a commercial service for language and grammar.

Reply: Thanks so much for your suggestions. We have a native English-speaking colleague edited the language of our manuscript. I hope that the current version is ready for publishing.

2) In some parts the manuscript is also quite demanding for readers, who are not specialist to the field, so a bit more explanation on experimental systems in the results section would be helpful for the general readership.

Reply: We thank Reviewer for pointing this out to us. We have explained our rationale for designing our experiments before going into results. We hope that our current version is more friendly to broader readership.

3) Especially the model system of tumor-SN-treated Raw264.7 needs a (short) introduction.
**Reply:** We thank Reviewer for bringing this point to our attention. We have now added a short introduction of the model system of tumor-SN-treated Raw264.7 (page9, line196-198).

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Overall, this manuscript by Zhou et al is interesting and timely. The authors identified a way to reprogram immunosuppressive M2 macrophages toward antitumor M1 cells, which may help to develop a novel immunotherapeutic strategy. The experiments are in general informative and reasonably well designed. However, several concerns on the analysis, presentation and interpretation of data are raised (see below).

**Reply:** We thank Reviewer for the nice comments on our study.

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In this manuscript, Zhou et al found that carfilzomib, a proteasome inhibitor (PI), could drive murine and human M2 macrophages to partially exhibit M1-like phenotype and function. By screening FDA-approved drugs, the authors identified three PIs namely carfilzomib, bortezomib and MLN9708 that effectively elicited IL-1β expression in murine bone marrow-derived macrophages (BMDMs). Further examination revealed that carfilzomib-induced M1-like cells (Ci-M1). Importantly, Ci-M1 exhibited enhanced phagocytotic and antigen-presenting activity, suggesting that carfilzomib treatment at least partially endowed M2/TAM with phenotype and functions similar to M1 macrophages. Investigation into the underlying mechanism of carfilzomib-induced M2 transformation into M1-like cells indicated that this process was dependent on an IRE1α-TRAF2-NF-κB pathway. Therapeutically, systemic carfilzomib treatment could inhibit tumor growth and synergize with PD-1 inhibitors in vivo, an effect that was attenuated in the absence of macrophages.

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1) Figure 1B: Three red dots are shown but they are not individually and specifically annotated. In addition, what does the x axis stand for? It is also unclear whether carfilzomib, bortezomib and MLN9708 are the only drugs in the library that were capable of activating IL-1β expression in IL4-induced M2 macrophages. Is there other PIs in the library that were not able to induce IL-1β expression? These issues should be clarified.

Reply: Following Reviewer’s suggestion, we have now annotated these three dots separately (New Figure 1B). We are sorry to make Reviewer confused. The X axis stands for the code of the drugs. We have now added the information in figure legend (Page 34, Line 950).

We checked our library carefully again. We confirmed that there are only three PIs in our library. During our screening, we found only Carfilzomib, Bortezomib and MLN9708 could activate IL-1β expression in IL4-induced M2 macrophages. We have now clarified this issue in the text (page 9, line 212).

2) Figure 2A: Although carfilzomib could decrease CD206 and arginase 1 expression, this reduction was marginal. Are the levels of IL-6 and iNOS expression post carfilzomib treatment comparable to that in conventional M1 macrophages? A positive control is recommended for supporting the authors’ claim that carfilzomib could reprogram M2 macrophage into M1-like population.

Reply: Following the reviewers’ suggestion, we compared the levels of Il-6 and Inos in Ci-M1 and conventional M1 macrophages (Figure 2a for Reviewer). The results indicated that Ci-M1 expressed lower levels of Il-6 and Inos compare with conventional M1 macrophages (LPS induced macrophage).

Chloroquine (CQ), a proven anti-malarial drug, could switch TAMs from M2 to M1 phenotype (Nature Communications (2018) 9: 873). Following the reviewers’
suggestion, we confirmed that CQ and Carfilzomib could reprogram M2 macrophage into M1-like population (Figure 2b for Reviewer). Both CQ and Carfilzomib can promote the expression of *Il-6* and *Inos*, Carfilzomib being more potent.

**Figure 2a for Reviewer**

![Figure 2a](image)

**Figure 2b for Reviewer**

![Figure 2b](image)

**Figure 2 for Reviewer:** (a) BMDMs/Raw264.7 cells were treated by LPS to transform into conventional M1 macrophages or IL-4 to transform into M2 macrophages, then Carfilzomib was added into IL-4 induced cells to further induce the formation of Ci-M1. *Il-6* and *Inos* were checked by qRT-PCR. (b) BMDMs/Raw264.7 cells were induced with IL-4 to transform into M2 macrophages, then treated by CQ or Carfilzomib. *Il-6* and *Inos* were checked by qRT-PCR.

3) Figure 2C: The gating strategy is unclear. How was the CD86+ and CD206+ population defined? Was it based on isotype-matched antibody staining or fluorescence minus one (FMO) control? This point is also applicable for other flow cytometric plots shown in the manuscript (e.g., Figure S2E and S2L).

**Reply:** Thanks very much for reminding us this important point. We defined the CD86+, CD206+, MHC-I+ and MHC-II+ population according to isotype-matched antibody staining control. We have added this important point in page 39, line...
4) Figure 2F and 2H: Both assays showed that carfilzomib could enhance macrophage phagocytotic ability, but the phagocytosis efficiencies exhibit a ten-fold difference. This should be clarified.

Reply: We thank Reviewer for pointing this out to us. Figure 2F showed that the number of L1210-GFP cells were phagocytosed per 100 macrophages. In some cases, one macrophage may phagocytose multiple L1210 cells. Figure 2H indicated that the percentage of phagocytosing macrophage in total macrophage. We are using these different methods for describing phagocytosing efficiency induced by Carfilzomib.

5) Figure 2L, y axis: Is it CD8+ % or CD4+ %? Figure 3D: Extra asterisks are shown on IL-6.

Reply: Reviewer is correct. The Y axis is CD4+ % in Figure 2L (new Figure 2L). Following Reviewer’s suggestion, we have now removed the extra asterisks in new Figure 3D.

6) Figure 4: It is very interesting that the kinase activity of IRE1α, instead of the endoribonuclease activity and the activation of XBP1, mediated the carfilzomib induced reprograming of M2 macrophages. The authors are encouraged to discuss the impact of PIs and ER stress inducers on XBP1-deficient cells.

Reply: Thanks for this great comment. Our results indicated that Kira6 (inhibiting IRE1 kinase activities), but not 4μ8C (inhibiting IRE1 RNase activities), inhibited Il-1β luciferase activities in Il-1β-luciferase transgenic Ci-M1 (Figure 4C & D). Stimulation of IRE1α activates both its kinase activity (to recruit TRAF2) and endoribonuclease activity (to splice XBP1). Spliced XBP1 transcribes genes involved in lowering unfolded protein load, including EIF2, AK3, Hspa5 and DNAJB9 (Genome Med. 2018 Oct 24;10(1):76). However, Kinase activity of IRE1α recruited TRAF2 to activate NF-xB to transcribe genes encoding M1 marker genes. We have
discuss it in Page 16-17, line 413-431.
Thank you for the submission of your revised manuscript to EMBO Molecular Medicine. We have now received the enclosed report from the two referees who were asked to re-assess it. As you will see the referees are now supportive, and I am pleased to inform you that we will be able to accept your manuscript pending the following amendments:

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2. Appendix
   - The file of merged Appendix Tables needs to be named as "Appendix".
   - Please add a table of contents to the first page of the appendix PDF file.

3. EV figures:
   - As we accept only up to 5 EV figures, please move at least 2 EV figures to the Appendix pdf, using the nomenclature Appendix Figure S1, etc, and please move their legends to the Appendix file.
   - Please also update the callouts in the main article, and make sure that all Appendix figures are called for.
   - Please upload EV figures as individual, high-resolution figures.

4. Source data: please note that source data needs to be directly linked to specific figures.
   - Source data for main figures: should be uploaded as one (zipped) file /figure, and named as "manuscriptID_SourceDataForFigure x"
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I look forward to seeing a revised version of your manuscript as soon as possible.

Kind regards
Jingyi
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2) Separate figure files*

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   - the medical issue you are addressing,
   - the results obtained and
   - their clinical impact.
   This may be edited to ensure that readers understand the significance and context of the research. Please refer to any of our published articles for an example.

6) For more information: There is space at the end of each article to list relevant web links for further consultation by our readers. Could you identify some relevant ones and provide such information as well? Some examples are patient associations, relevant databases, OMIM/proteins/genes links, author's websites, etc...

7) Author contributions: the contribution of every author must be detailed in a separate section.

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9) Every published paper now includes a 'Synopsis' to further enhance discoverability. Synopses are displayed on the journal webpage and are freely accessible to all readers. They include a short stand first (maximum of 300 characters, including space) as well as 2-5 one sentence bullet points that summarise the paper. Please write the bullet points to summarise the key NEW findings. They should be designed to be complementary to the abstract - i.e. not repeat the same text. We encourage inclusion of key acronyms and quantitative information (maximum of 30 words / bullet point). Please use the passive voice. Please attach these in a separate file or send them by email, we will incorporate them accordingly.
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***** Reviewer's comments *****

Referee #1 (Comments on Novelty/Model System for Author):
The topic is important and closely related to the translational research.

Referee #1 (Remarks for Author):
The authors have addressed my concerns and the revised version is improved. I have no further questions.

Referee #3 (Remarks for Author):
The authors have addressed all the concerns raised by the Reviewers and have improved the manuscript accordingly.
The authors performed the requested editorial changes.
5th Nov 2021

Dear Dr. Chen,

Please find enclosed the final reports on your manuscript. We are pleased to inform you that your manuscript is accepted for publication and is now being sent to our publisher to be included in the next available issue of EMBO Molecular Medicine.

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Jingyi Hou
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A- Figures

1. Data

The data shown in figures should satisfy the following conditions:

- The data were obtained and processed according to the field's best practice and are presented to reflect the results of the experiments in an accurate and unbiased manner.
- Figure panels include only data points, measurements or observations that can be compared to each other in a scientifically meaningful way.
- Graphs include clearly labeled error bars for independent experiments and sample sizes. Unless justified, error bars should not be shown for technical replicates.
- If n=5, the individual data points from each experiment should be plotted and any statistical test employed should be justified.
- Source Data should be included to report the data underlying graphs. Please follow the guidelines set out in the authorship guidelines on Data Presentation.

2. Captions

Each figure caption should contain the following information, for each panel where they are relevant:

- A description of the sample collection allowing the reader to understand whether the samples represent technical or biological replicates (including how many animals, litters, cultures, etc.).
- A description of the sample collection allowing the reader to understand whether the samples represent technical or biological replicates (including how many animals, litters, cultures, etc.).
- A specification of the experimental system investigated (e.g. cell line, species name).
- An explicit mention of the biological and chemical entity(ies) that are altered/varied/perturbed in a controlled manner.
- An explicit mention of the biological and chemical entity(ies) that are being measured.
- A statement of how many times the experiment shown was independently replicated in the laboratory.
- Definitions of statistical methods and measures:
  - A statement of how many times the experiment shown was independently replicated in the laboratory.
  - Common tests, such as t-test (please specify whether paired or unpaired), simple χ2 tests, Wilcoxon and Mann-Whitney tests, can be unambiguously identified by name only, but more complex techniques should be described in the methods section.
  - Are tests one-sided or two-sided?
  - Are there adjustments for multiple comparisons?
  - Exact statistical test results, e.g., P values = x but not P values < x;
  - Definition of ‘center values’ as median or average;
  - Definition of error bars as s.d. or s.e.m.

Any descriptions too long for the figure legend should be included in the methods section and/or with the source data.

In the pink boxes below, please ensure that the answers to the following questions are reported in the manuscript itself. Every question should be answered. If the question is not relevant to your research, please write NA (non applicable). We encourage you to include a specific subsection in the methods section for statistics, reagents, animal models and human subjects.

If the pink boxes below, please ensure that the answers to the following questions are reported in the manuscript itself. Every question should be answered. If the question is not relevant to your research, please write NA (non applicable). We encourage you to include a specific subsection in the methods section for statistics, reagents, animal models and human subjects.

B- Statistics and general methods

| Question                                                                 | Answer                          |
|-------------------------------------------------------------------------|---------------------------------|
| 1a. How was the sample size chosen to ensure adequate power to detect a pre-specified effect size? | Samples size was set such that statistical significance can be reached. |
| 1b. For animal studies, include a statement about sample size estimate even if no statistical methods were used. | Statistical analysis was done on all experiments that used mice. We have indicated the mouse numbers in Materials and Methods and in Figure legends. |
| 2. Describe inclusion/exclusion criteria if samples or animals were excluded from the analysis. Were the criteria pre-established? | In our experiment when we used Cal-97 tumors or other treatments (as indicated in the main text) to treat transgenic mice, computed tomography was conducted to confirm lung cancer in mice. Mice bearing no noticeable lung cancers before treatment were excluded from analysis. |
| 3. Were any steps taken to minimize the effects of subjective bias when allocating animals/samples to treatment (e.g., randomization procedure)? If yes, please describe. | In our treatment studies, we chose a cohort of mice bearing similar burden of lung cancer. These mice were then randomized to different treatments. |
| 4a. Were any steps taken to minimize the effects of subjective bias during group allocation or/and when assessing results (e.g., blinding of the investigator)? If yes please describe. | The mice were randomized for treatment. We have indicated this in main text. |
| 4b. For animal studies, include a statement about blinding even if no blinding was done. | Blinding software was used to analyze the CT images for determining tumor burdens. |
| 5. For every figure, are statistical tests justified as appropriate? | Yes, statistical tests were justified as appropriate. |
| Do the data meet the assumptions of the tests (e.g., normal distribution)? Describe any methods used to assess it. | Not applicable |
| Is there an estimate of variation within each group of data? | No, there was no estimate of variation within each group of data. |
### C- Reagents

| Question                                                                 | Answer                                                                 |
|-------------------------------------------------------------------------|------------------------------------------------------------------------|
| Do antibodies used in your study fall under dual use research restrictions? | Yes, this information is stated in Materials and Methods. All animals were housed in specific pathogen-free conditions and breeding and all animal procedures were conducted in strict accordance with guidelines for the care and use of laboratory approved by the Institute of Laboratory Animal Science, Jinan University. |