Succession of Transiently Active Tumor-Initiating Cell Clones in Human Pancreatic Cancer Xenografts

Claudia R. Ball, Felix Oppel, K. Roland Ehrenberg, Taronish D. Dubash, Sebastian M. Dieter, Christopher M. Hoffmann, Ulrich Abel, Friederike Herbst, Moritz Koch, Jens Werner, Frank Bergmann, Naveed Ishaque, Manfred Schmidt, Christof von Kalle, Claudia Scholl, Stefan Fröhling, Benedikt Brors, Wilko Weichert, Jürgen Weitz, Hanno Glimm

Corresponding author: Hanno Glimm, National Center for Tumor Diseases NCT

Review timeline:

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Transaction Report:

(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. The original formatting of letters and referee reports may not be reflected in this compilation.)

Editor: Roberto Buccione

1st Editorial Decision 21 December 2016

Thank you for the submission of your manuscript to EMBO Molecular Medicine.

I apologize for the delay in providing you with a decision. We experienced significant difficulties in securing willing and appropriate reviewers and then obtaining their evaluations in a timely fashion. Also, I have not been able to obtain a third evaluation on this manuscript. Hence I have decided to proceed based on the two available evaluations to avoid further delays.

As you will see, while reviewer 2 appears more positive, reviewer 1 is rather more reserved. Specifically, reviewer 1’s main concern is the transplantation procedure and a number of parameters related to this, heavily affect the outcome and consequent interpretation of the results and does not feel that the current dataset actually informs us on hierarchical and clonal architecture. Reviewer 2, while less reserved, does ask two poignant questions, complementary and partially overlapping with reviewer 1’s concerns: the relationship between clonal dynamics and metastatic growth and the fact that the limited number of tumors considered might not allow for generalization.

After further cross-commenting with the reviewers it was agreed that the study is potentially very interesting but it is currently lacking in conclusiveness and sufficient experimental support for the main claims.

In conclusion, I am prepared to consider a substantially revised submission, with the understanding
that the Reviewers' concerns must be addressed, especially to verify the relevance of your findings by comparing clonal representation in primary tumors and metastasis, since metastasis may be a physiological process that somewhat recapitulates an experimental tumor initiation setting. As for the other limitations of the study, these should be at a minimum, properly acknowledged and appropriately discussed.

Please note that it is EMBO Molecular Medicine policy to allow a single round of revision only and that, therefore, acceptance or rejection of the manuscript will depend on the completeness of your responses included in the next, final version of the manuscript.

As you know, 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. However, I do ask you to get in touch with us after three months if you have not completed your revision, to update us on the status. Please also contact us as soon as possible if similar work is published elsewhere.

Please note that EMBO Molecular Medicine now requires a complete author checklist (http://embomolmed.embopress.org/authorguide#editorial3) to be submitted with all revised manuscripts. Provision of the author checklist is mandatory at revision stage; The checklist is designed to enhance and standardize reporting of key information in research papers and to support reanalysis and repetition of experiments by the community. The list covers key information for figure panels and captions and focuses on statistics, the reporting of reagents, animal models and human subject-derived data, as well as guidance to optimise data accessibility.

We now mandate that all corresponding authors list an ORCID digital identifier. You may acquire one through our web platform upon submission and the procedure takes <90 seconds to complete. We also encourage co-authors to supply an ORCID identifier, which will be linked to their name for unambiguous name identification.

I look forward to seeing a revised form of your manuscript as soon as possible.

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

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

The model system is suitable to study tumor initiation in immune compromised mice but has no relevance for established cancer tissue as is claimed.

Referee #1 (Remarks):

With interest I read the manuscript "Succession of transiently active tumor-initiating cell clones in human pancreatic cancer" by Claudia R. Ball et al.

The authors clonally marked primary human pancreatic ductal adenocarcinoma (PDAC) cells by lentiviral transduction and serially transplanted these xenografts. With this model the authors aim to monitor the clonal dynamics and self-renewal activity of clones during tumor formation in vivo. Rather surprisingly, by analyzing the different integration sites of the vector, using LAM-PCR, they found that almost no overlap in integration sites between the cells of the primary, secondary and tertiary xenografts was present. The authors conclude this indicates distinct transiently active populations of TICs are present, generating xenograft tumors after transplantation in PDAC xenografts.

The manuscript is well written and the experiments well described and have been executed to the highest standards. However, I have some serious doubts about the interpretation of the findings.

Major comments/recommendations
- The most important criticism involves the transplantation procedure itself. It is concluded that the continuous appearance of 'new' clones is due to differential activation of various TIC populations, and potential exhaustion of previous highly proliferative clones.

Firstly, the cells that are analyzed for their integration side are by nature of the procedure not the
cells that are serially transplanted. Hence, there is an important selection taking place here. Secondly, and more importantly, the stochasticity of the seeding determines the eventual results of the clonal outgrowth. Apparently only a very limited number of cells after injection substantially contributes to each xenograft, and a large proportion of cells simply sit there because they are on the inside of the tumour(?) and are quickly outgrown by cells on the outside of the tumour. This has nothing to do with stem cell potential.

- In relation to the point above, even if based on the current data conclusions can be drawn on stem cell function in cancer this only involved the initiation phase of xenografts not the dynamics of stem cells in established cancer tissue. This is an important drawback.

- It would be interesting to see if a difference in proliferation rate can be shown in the xenograft tumors and if there are specific regions where there is more or less proliferation. Now solely based on mathematical analysis of the data, this proliferation difference was explained. However, this was not clearly evident from the experiments; could the researchers understated their findings with an experiment (Ki67 staining)? This could possibly strengthen their findings.

- Did the authors find metastases of the renal capsule and orthotopic PDAC xenograft transplantations? According to their previous paper (Dieter et al, 2011), where they examined the TIC in human colon cancer, they showed that metastasis formation was almost exclusively driven by self-renewing long-term-TICs. It would be interesting to see which population of TIC is driving metastases in PDAC as well.

- I think the plural TICs should be used at many instances instead of TIC.

Referee #2 (Remarks):

"Succession of Transiently Active Tumor-Initiating Cell Clones in Human Pancreatic Cancer"

Previous studies had identified populations of tumor initiating cells in pancreas tumors with the capacity to propagate the disease in mice. It was thus proposed that pancreas cancer follows a hierarchical organization similar to that present in other cancer types. These assays, however, were largely biased by the choice of surface markers used to identify and isolate tumor cell populations. In addition, tumor initiation assays only provide information about the behavior of isolated tumor cells. In this manuscript, Glimm and colleagues used lentiviral marking of primary pancreatic cancer cultures to study tumor growth. In many aspects, this approach is superior to transplantation assays as it enables an unbiased assessment of clone dynamics on whole tumors. Authors had previously applied a similar approach to study Colorectal Cancer (CRC) where they reached the conclusion that CRCs are maintained by a hierarchy of cells that includes cancer stem cells at the apex. Therefore, clonal dynamics assessed through lentiviral marking validated the model that had been previously elaborated from tumor cell transplantation assays in CRC. Using an equivalent approach, authors now reach the surprising conclusion that Pancreatic cancer is not organized according to a hierarchy but rather that long-term growth in this type of tumors is maintained by successive recruitment of inactive clones that are activated in restricted periods. Fittingly, authors also demonstrate that frequencies of TICs in ex vivo cultures do not correlate with expression of stem or differentiation markers. Altogether, these are unexpected observations with profound implications to understand pancreatic cancer and improve therapeutic treatments. In my opinion, data are convincing and of high quality. I only have minor criticism/suggestions:

- Does metastatic growth follow clonal dynamics similar to the observed for primary xenografts? This is an important issue as pancreatic cancer is highly metastatic and therapies are not effective to treat metastasis.

- Authors explore the behavior of a small number of primary tumors and therefore it is possible that author's conclusions only hold for a subset of pancreatic cancers whereas other types may follow a hierarchical organization. Authors should remain open to this possibility in the discussion.
TICs have been described to drive development and progression of a variety of human cancers, including pancreatic cancer. We agree with the Referee that the functional capacity of human TIC cannot be analyzed within the natural undisturbed in vivo situation, i.e. within the patient. Instead, experimental analysis of TIC biology in humans by nature requires surgical removal of cancer tissue, dissociation of the patient tumor and subsequent functional readouts in adequate in vivo and in vitro surrogate models. Still, by adapting functional assays originally developed for normal adult stem cells, key properties of TIC have been successfully investigated (O’Brien et al Nature 2007, Ricci-Vitiani et al Nature 2007; Dalerba et al, PNAS 2007; Li et al, Cancer Res 2007; Hermann et al. Cell Stem Cell 2007). In this context, serial transplantation of human cancer cells into immunodeficient mice has been widely used to quantify self-renewal of TIC and tumor long-term progression. We adapted these well-established approaches within our study and unexpectedly demonstrate that PDAC progression in this model unlike colon cancer does not require stably self-renewing tumor stem cells within a fixed malignant stem cell compartment. We believe that these findings are highly relevant for the development of therapeutic approaches targeting TIC activity in pancreatic cancer. We added a paragraph to discuss the limitations of the experimental model used.

Referee #1 (Remarks):
With interest I read the manuscript "Succession of transiently active tumor-initiating cell clones in human pancreatic cancer" by Claudia R. Ball et al.
The authors clonally marked primary human pancreatic ductal adenocarcinoma (PDAC) cells by lentiviral transduction and serially transplanted these xenografts. With this model the authors aim to monitor the clonal dynamics and self-renewal activity of clones during tumor formation in vivo. Rather surprisingly, by analyzing the different integration sites of the vector, using LAM-PCR, they found that almost no overlap in integration sites between the cells of the primary, secondary and tertiary xenografts was present. The authors conclude this indicates distinct transiently active populations of TICs are present, generating xenograft tumors after transplantation in PDAC xenografts.

The manuscript is well written and the experiments well described and have been executed to the highest standards.

We thank the Reviewer for his appreciation of our experimental work.

However, I have some serious doubts about the interpretation of the findings.

Major comments/recommendations
1. The most important criticism involves the transplantation procedure itself. It is concluded that the continuous appearance of ’new’ clones is due to differential activation of various TIC populations, and potential exhaustion of previous highly proliferative clones.
Firstly, the cells that are analyzed for their integration side are by nature of the procedure not the cells that are serially transplanted. Hence, there is an important selection taking place here. Secondly, and more importantly, the stochasticity of the seeding determines the eventual results of the clonal outgrowth. Apparently only a very limited number of cells after injection substantially contributes to each xenograft, and a large proportion of cells simply sit there because they are on the inside of the tumour(?) and are quickly outgrown by cells on the outside of the tumour. This has nothing to do with stem cell potential.

Testing engraftment and cellular progeny generation in serial transplantation is a long-proven standard approach for assaying stem cell activity within a given cell population. In combination with clonal marking, this strategy has been conclusively shown to enable tracing clonal activity of individual normal and benign stem cells in vivo. Of note, using the same experimental approach as described here, we recently characterized the organization of the tumor-initiating cell (TIC) compartment in human colon cancer and came to fundamentally different results. Our highly sensitive technology was able to precisely determine the clonal contribution within a hierarchically organized TIC compartment with self-renewing TIC that give rise to transient tumor-amplifying cells in colon cancer xenografts (Dieter, S.M., et al., Cell Stem Cell, 2011. 9(4): p. 357-65).

In the study presented here, the majority of marked PDAC cell clones that strongly contributed to tumor formation in one generation were not detectable in other serial xenograft generations. Importantly, at each step of serial transplantation, single cell suspensions have been generated and injected, a strategy that ensures equal distribution of cell clones within the transplant. Nevertheless, the majority of cell clones that contributed strongly to tumor formation
in one generation were not detected in subsequent generations. As within a given tumor generation, more than 90% of all marked cells descended from a set of tumor-specific dominant clones and thereby represented the vast majority of all cells which are transplanted into subsequent mice this is a highly surprising and unexpected finding. Although these cells outnumber all others by far at any given location they did not contribute to tumor-formation in subsequent xenografts after serial transplantation to a measurable extent by our highly sensitive LAM PCR. This cannot be explained by a stochastic contribution of individual cancer cells but indicate that clonal output was transient and did not depend on intra-tumor localization. In line with this, Ki67 staining of xenografts that we now added as suggested by the reviewer clearly shows that proliferating cancer cells are equally distributed throughout the tumors. In addition, our in vitro data demonstrate similar clonal dynamics in serial cultures that by nature cannot be explained by the localization of cells within tumors. We adjusted the manuscript to make these points clearer.

2. In relation to the point above, even if based on the current data conclusions can be drawn on stem cell function in cancer this only involved the initiation phase of xenografts not the dynamics of stem cells in established cancer tissue. This is an important drawback.

We agree that serial transplantation of human cells in a xenograft model requires disruption of the established cancer tissue by singularization of cells and subsequent injection into the xenogeneic graft. However, despite these limitations, so far this is the only reliable assay to monitor human (tumor) stem cell activity in vivo as dynamics of human stem cells cannot be assayed directly in patients. We would like to point out that serial transplantation of clonally marked cells is well established in the field of stem cell research to determine the self-renewal capacity and long-term growth of initially transplanted cell populations (please see also answer to point 1 raised by the same referee). As stated above, we added a discussion of the limitations of the experimental model used to the revised version of the manuscript.

3. It would be interesting to see if a difference in proliferation rate can be shown in the xenograft tumors and if there are specific regions where there is more or less proliferation. Now solely based on mathematical analysis of the data, this proliferation difference was explained. However, this was not clearly evident from the experiments; could the researchers understate their findings with an experiment (Ki67 staining)? This could possibly strengthen their findings.

We thank the Referee for this valuable suggestion. We now have performed Ki67 staining of mouse PDAC xenografts as requested. These data clearly demonstrate the presence of cells with different proliferative activities, i.e. actively proliferating and inactive cells. Strikingly, proliferating cells are equally distributed throughout the tumor, showing that active proliferation is present within the whole tumor mass and not restricted to specific sites of the tumor, i.e. the outside as suggested in comment 1 of the same Referee. As discussed in our response to point 1 these results strongly support our conclusion that the differences in proliferative activity of clones in serial transplantation cannot simply be explained by their spatial distribution, especially as dominant TIC clones make up the vast majority of cells at any given location within the tumor and thereby also supply the vast majority of cells re-transplanted.

4. Did the authors find metastases of the renal capsule and orthotopic PDAC xenograft transplantations? According to their previous paper (Dieter et al, 2011), where they examined the TIC in human colon cancer, they showed that metastasis formation was almost exclusively driven by self-renewing long-term-TICs. It would be interesting to see which population of TIC is driving metastases in PDAC as well.

We agree that understanding the clonal dynamics of metastasis formation would be highly interesting. However, unlike our colon cancer models, the human pancreatic cancer xenografts do not metastasize spontaneously. Accordingly, clonal analysis can only be done on PDAC xenograft tumors and clonal analysis of metastasis formation is simply not possible. Of note, a potential development of metastasis formation surrogate models, subsequent serial transplantation and molecular analyses if successful would require several additional years of experimentation in rather artificial models. We therefore feel that such analyses if at all feasible are –although highly interesting– beyond the scope of the current manuscript and need to be addressed separately.

5. I think the plural TICs should be used at many instances instead of TIC.

We agree and changed the manuscript accordingly.
Referee #2 (Remarks):

"Succession of Transiently Active Tumor-Initiating Cell Clones in Human Pancreatic Cancer"

Previous studies had identified populations of tumor initiating cells in pancreas tumors with the capacity to propagate the disease in mice. It was thus proposed that pancreas cancer follows a hierarchical organization similar to that present in other cancer types. These assays, however, were largely biased by the choice of surface markers used to identify and isolate tumor cell populations. In addition, tumor initiation assays only provide information about the behavior of isolated tumor cells. In this manuscript, Glimm and colleagues used lentiviral marking of primary pancreatic cancer cultures to study tumor growth. In many aspects, this approach is superior to transplantation assays as it enables an unbiased assessment of clone dynamics on whole tumors. Authors had previously applied a similar approach to study Colorectal Cancer (CRC) where they reached the conclusion that CRCs are maintained by a hierarchy of cells that includes cancer stem cells at the apex. Therefore, clonal dynamics assessed through lentiviral marking validated the model that had been previously elaborated from tumor cell transplantation assays in CRC. Using an equivalent approach, authors now reach the surprising conclusion that Pancreatic cancer is not organized according to a hierarchy but rather that long-term growth in this type of tumors is maintained by successive recruitment of inactive clones that are activated in restricted periods. Fittingly, authors also demonstrate that frequencies of TICs in ex vivo cultures do not correlate with expression of stem or differentiation markers. Altogether, these are unexpected observations with profound implications to understand pancreatic cancer and improve therapeutic treatments.

We thank the Reviewer for his precise synopsis.

In my opinion, data are convincing and of high quality. I only have minor criticism/suggestions:

- Does metastatic growth follow clonal dynamics similar to the observed for primary xenografts? This is an important issue as pancreatic cancer is highly metastatic and therapies are not effective to treat metastasis.

We agree that understanding the clonal dynamics of metastasis formation would be highly interesting. However, unlike our colon cancer models, the human pancreatic cancer xenografts do not metastasize spontaneously. Accordingly, clonal analysis can only be done on PDAC xenograft tumors and clonal analysis of metastasis formation is simply not possible. Of note, a potential development of metastasis formation surrogate models, subsequent serial transplantation and molecular analyses if successful would require several additional years of experimentation in rather artificial models. We therefore feel that such analyses if at all feasible are –although highly interesting- beyond the scope of the current manuscript and need to be addressed separately.

- Authors explore the behavior of a small number of primary tumors and therefore it is possible that author's conclusions only hold for a subset of pancreatic cancers whereas other types may follow a hierarchical organization. Authors should remain open to this possibility in the discussion.

We agree with the Reviewer and adjusted the discussion accordingly.

2nd Editorial Decision 29 March 2017

Thank you for the submission of your revised manuscript to EMBO Molecular Medicine. We have now received the enclosed reports from the reviewers that were asked to re-assess it. As you see, while reviewer 2 is now supportive, reviewer 1 is decidedly more reserved.

Reviewer 1 is not satisfied that your revision addresses his/her main concern that your approach is informative regarding the properties of growing cancers and therefore that ultimately the conclusions are not adequately supported by the data.

After further discussion, reviewer 1 reiterated his/her position but also appreciation for an interesting, technically well-performed set of experiments. The reviewer also acknowledges that s/he has a different interpretation of your data. Ultimately however, s/he would not be opposed to publication of your manuscript, provided the findings were better contextualized and the limitations more clearly stated. I agree and also suggest that the spirit of reviewer 2's comments would also be best served by such a revision.
I am pleased to inform you that we will be able to accept your manuscript, provided you carefully consider and act upon the above. Please make sure the changes are highlighted in the manuscript. Finally, please also note the following pending final amendments:

1) We note that Fig S2 is called out in the manuscript before Fig. S1. Please reorganize in chronological order. We also note that there are no callouts for Tables S5, S6.

2) As per our Author Guidelines, the description of all reported data that includes statistical testing must state the name of the statistical test used to generate error bars and P values, the number (n) of independent experiments underlying each data point (not replicate measures of one sample), and the actual P value for each test (not merely 'significant' or 'P < 0.05').

3) The manuscript must include a statement in the Materials and Methods identifying the institutional and/or licensing committee approving the experiments, including any relevant details (like how many animals were used, of which gender, at what age, which strains, if genetically modified, on which background, housing details, etc). We encourage authors to follow the ARRIVE guidelines for reporting studies involving animals. Please see the EQUATOR website for details: http://www.equator-network.org/reporting-guidelines/improving-bioscience-research-reporting-the-arrive-guidelines-for-reporting-animal-research/. Please make sure that ALL the above details are reported, including in the checklist.

4) We encourage the publication of source data, with the aim of making primary data more accessible and transparent to the reader. Would you be willing to provide a PDF file per figure that contains the original, uncropped and unprocessed scans of all or at least the key gels used in the manuscript and/or source data sets for relevant graphs? The files should be labeled with the appropriate figure/panel number, and in the case of gels, should have molecular weight markers; further annotation may be useful but is not essential. The files will be published online with the article as supplementary "Source Data" files. If you have any questions regarding this just contact me.

5) We encourage the provision of striking image or visual abstract to illustrate your article. If you do, please provide a jpeg file 550 px-wide x 400 px high.

Please submit your revised manuscript within two weeks. I look forward to seeing a revised form of your manuscript as soon as possible.

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

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

The model system solely allows for the study of tumor initiation, and is not informative regarding properties of growing cancers. The conclusions of the authors are not supported by the data.

Referee #1 (Remarks):

As I also indicated in my initial review in my view there is an important conceptual problem with the model. All of the effects that are reported are potential artefacts of the engraftment phase of the assay and not of the features of tumor growth. Therefore, the conclusions as drawn by the authors, are not supported by the data.

The reply of the authors to my concerns is not very convincing, and does not present any new insights. Indeed, ten years ago the tumor initiation assay was accepted to be related to stem cells. In 2017 better assays are required to make claims about potential distinct types of stem cells within malignancies. Unfortunately, no new data could be presented on natural engraftment, i.e. metastasis formation, as was also requested by the other reviewer.
Referee #2 (Remarks):

Authors argue that pancreatic model systems that they use do not generate metastasis, which precludes performing the experiments that I suggested. Instead, authors have discussed the limitations of the system in this revised version. My impression is that the work is relevant and should be published. Despite the technical caveats and potential artifacts that we discussed with the other reviewer (and that I subscribe), it is important to consider that similar transplantation experiments have been widely used to assess the hierarchical organization of other tumor types. In the case of CRC, authors applied the same lentiviral marking followed by serial transplantation. They observed, however, that CRC is hierarchically organized, which argues against the possibility that their conclusion in pancreas cancer about clonal succession may simply represent a byproduct of the transplantation assay.

2nd Revision - authors' response 11 April 2017

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

The model system solely allows for the study of tumor initiation, and is not informative regarding properties of growing cancers. The conclusions of the authors are not supported by the data.

Xenotransplantation of human cancer cells into severely immune deficient mice is widely used due to the lack of alternate experimental in vivo models for human cancer. We agree that it cannot be excluded that disruption of the original tumor’s architecture, the transplantation procedure and the xenogenic environment may influence the behavior of the assayed tumor cell population. However, these assays have been adopted to successfully describe the presence and key properties, i.e. self-renewal and long-term activity, of tumor initiating cells in a wide range of human cancer tissue. We adapted these well-established approaches within our study and unexpectedly demonstrate that PDAC serial transplantation in this model unlike colon cancer does not require stably self-renewing tumor stem cells within a fixed malignant stem cell compartment. In the revised version of the manuscript, we have expanded the discussion of the limitations of the experimental model used and clearly contextualized the obtained results.

Referee #1 (Remarks):

As I also indicated in my initial review in my view there is an important conceptual problem with the model. All of the effects that are reported are potential artefacts of the engraftment phase of the assay and not of the features of tumor growth. Therefore, the conclusions as drawn by the authors, are not supported by the data. The reply of the authors to my concerns is not very convincing, and does not present any new insights. Indeed, ten years ago the tumor initiation assay was accepted to be related to stem cells. In 2017 better assays are required to make claims about potential distinct types of stem cells within malignancies. Unfortunately, no new data could be presented on natural engraftment, i.e. metastasis formation, as was also requested by the other reviewer.

Indeed, continuous tumor growth cannot be assayed in serial xenotransplantation, as tumor explantation and dissociation before xenotransplantation are required. Therefore, it cannot be ruled out that the procedure of xenotransplantation per se impacts the clonal dynamics observed. However, we would like to emphasize that by using the same methodology we have previously demonstrated a hierarchical organization of the TIC compartment in colorectal cancer (Dieter et al., 2011), clearly establishing that the experimental model used is permissive for fundamentally different clonal dynamics of patient samples from different malignant diseases. Moreover, within the current study, we detected very similar clonal dynamics of cultured PDAC cells in vitro as we observed in serial xenografts in vivo, arguing against a dominant effect of xenogeneic engraftment on clonal TIC dynamics. Within the new version of the manuscript, we have thoroughly discussed the potential limitations of the model system in the introduction and discussion sections. Moreover, we have modified the title and text throughout the manuscript to more clearly indicate that the data presented are generated in the context of serial xenotransplantation.

Referee #2 (Remarks):

Authors argue that pancreatic model systems that they use do not generate metastasis, which precludes performing the experiments that I suggested. Instead, authors have discussed the limitations of the system in this revised version. My impression is that the work is relevant and should be published. Despite the technical caveats and potential artifacts
that we discussed with the other reviewer (and that I subscribe), it is important to consider that similar transplantation experiments have been widely used to assess the hierarchical organization of other tumor types. In the case of CRC, authors applied the same lentiviral marking followed by serial transplantation. They, observed, however, that CRC is hierarchically organized, which argues against the possibility that their conclusion in pancreas cancer about clonal succession may simply represent a byproduct of the transplantation assay.

We thank the reviewer for his positive comments.
Submitted to: EMBO Molecular Medicine
Manuscript Number: EMM-2016-07354-V3

Corresponding Author Name: Nanno Guillon

Reporting Checklist For Life Sciences Articles (Rev. July 2015)

This checklist is used to ensure good reporting standards and to improve the reproducibility of published results. These guidelines are consistent with the Principles and Guidelines for Reporting Preclinical Research issued by the NIH in 2014. Please follow the journal’s authority guidelines in preparing your manuscript.

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 specification of the experimental system investigated (e.g. cell line, species name).
- the assay(s) and method(s) used to carry out the reported observations and measurements.
- an explicit mention of the biological and chemical entity(ies) that are being measured.
- an explicit mention of the biological and chemical entity(ies) that are altered/changed/perturbed in a controlled manner.
- the exact sample size (n) for each experimental group/condition, given as a number, not a range.
- 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 statement of how many times the experiment shown was independently replicated in the laboratory.
- definitions of statistical methods and measures: common tests, such as t-test (please specify whether paired or unpaired), simple p-tests, Wilcoxon and Mann-Whitney tests, can be unambiguously identified by name only, but more complex techniques should be described in the methods section.
- tests one-sided or two-sided?
- are there adjustments for multiple comparisons?
- mean statistical test results, e.g., if p-value < 0.05 but not p-value < 0.01.
- 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.

Please ensure that the answers to the following questions are reported in the manuscript itself. 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

1.a. How was the sample size chosen to ensure adequate power to detect a pre-specified effect size?

1.b. For animal studies, include a statement about sample size estimate even if no statistical methods were used.

2 a. Describe inclusion/exclusion criteria of samples or animals were excluded from the analysis. Were the criteria pre-established?

2 b. Were any steps taken to reduce the effects of subjective bias when allocating animals/samples to treatment (e.g. randomization procedure)? If yes, please describe.

For animal studies, include a statement about randomization even if no randomization was used.

3 a. 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.

3 b. For animal studies, include a statement about blinding even if no blinding was done

4. For every figure, are statistical tests justified as appropriate?

5. Do the data meet the assumptions of the tests (e.g., normal distribution)? Describe any methods used to assess it.

6. Are there estimates of variation within each group of data?

7. Are the variances similar between the groups that are being statistically compared?

C- Reagents
D- Animal Models

1. Report species, strain, gender, age of animals and genetic modification status where applicable. Please detail housing and husbandry conditions and the source of animals.

2. For experiments involving live variables, include a statement of compliance with ethical regulations and identify the committee(s) approving the experiments.

3. We recommend considering the ARRIVE guidelines [see link list at top-right] (Cho et al., 2009, eLife00142, 2010) to ensure that other relevant aspects of animal studies are adequately reported. See author guidelines, under ‘Reporting Guidelines’. See also NIH [see link list at top-right] and MIRIAM guidelines [see link list at top-right] recommendations. Please confirm compliance.

4. Provide accession codes for deposited data. See author guidelines, under ‘Data deposition’.

5. Data deposition in a public repository is mandatory for:
   a. Proteins, DNA and RNA sequences
   b. Macromolecular structures
   c. Crystallographic data for small molecules
   d. Functional genomics data
   e. Proteomics and molecular interactions

6. If deposition is strongly recommended for any datasets that are central and integral to the study, please consider the journal’s data policy. If it is not structured public repository exists for a given data type, we encourage the provision of datasets in the manuscript as a Supplementary Document (see author guidelines under ‘Expanded View’ or in-institutional repositories such as Dryad [see link list at top-right] or Figshare [see link list at top-right]).

7. Access to human and animal datasets should be provided with clear instructions on how to access the data respecting ethical obligations to the patients and relevant medical and legal issues. If practically possible and compatible with the individual consent agreements used in the study, such data should be deposited in one of the major public access controlled repositories such as BioSample [see link list at top-right] or EGA [see link list at top-right].

8. Access to possible, primary and referenced data should be formally cited in a Data Availability section. Please state whether you have included this section.

   Examples:

   - Primary Data
     Referenced Data
     Huang J, Brown AF, Lei M (2012). Crystal structure of the TRBD domain of TRT and the OA1/5 of TR. Protein Data Bank 5036
     AF-MS analysis of human histone deacetylase interactions in CD4+ T cells (2011). PRIDE P0000018

   - Computational models that use animal and integral to a study should be shared without restrictions and provided in a machine-readable form. The relevant accession numbers or links should be provided. When possible, standardized formats (SBML, CellML) should be used instead of scripts (e.g., MATLAB). Authors are strongly encouraged to follow the MIRIAM guidelines [see link list at top-right] and deposit their model in a public database such as BioModels [see link list at top-right] or BioModels Online [see link list at top-right]. If a computer source code is provided with the paper, it should be deposited in a public repository or included in supplementary information.

G- Dual use research of concern

19. Could your study fall under dual use research restrictions? Please check biosecurity documents [see link list at top-right] and list of select agents and toxins [APHS/CDC] [see link list at top-right]. According to our biosecurity guidelines, provide a statement only if it could.