Chronic inflammation: is it the driver or is it paving the road for malignant transformation?

Kambiz Afrasiabi¹, Yi-Hong Zhou², Angela Fleischman¹

¹ Department of Medicine, University of California, Irvine, CA, USA
² Department of Surgery, University of California, Irvine, CA, USA

Correspondence to: Kambiz Afrasiabi, email: kambiz@uci.edu

Keywords: Chronic inflammation, Free energy, driver mutation, bioenergetics, metabolomics, electric field and cancer, JAK2⁶¹⁷⁵, p53 Rb

Received: March 17, 2015  Accepted: May 7, 2015  Published: May 8, 2015

This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

ABSTRACT

Chronic inflammation in well-defined mouse models such as Gia2 knock out mouse has been shown to trigger formation and expansion of hypoxic niches and also leads to up regulation of NFĸB, offering cells which have adapted their genetic machinery to hypoxia a unique survival advantage. These adapted cells have been shown to acquire stem cell-like capabilities as shown by up regulation of stem cell markers. Such long lived cells become permanent residents in sub mucosa and acquire a malignant phenotype from long-term exposure to noxious environmental agents due to a barrier defect secondary to down regulation of barrier proteins such as Zo1 and Occludin. Indeed mitotic spindle disorientation in such mice has been proposed as another contributory factor to malignant transformation. Sterilization of bowel lumen of these mice through different techniques has prevented malignant transformation in the presence of chronic inflammation. These facts stand strongly against chronic inflammation as a true driver of carcinogenesis but clearly support its role in facilitating the emergence of the neoplastic clone.

INTRODUCTION

The malignant transformation as per the free energy concept necessitates a significant and permanent decrease in the free energy of the malignant cell as a result of breakdown of the fine balance of cellular energetics[2]. Warburg effect offers a credible reflection of these deeply seated perturbances in cellular energetics and metabolism[4]. If chronic inflammation in well defined and valid mouse models would directly lead to break down of cellular energetics, then the assignment of a driver role to it would become meaningful. However evidence points to the contrary as discussed below. Indeed the cells that evolve in the inflammatory microenvironment under the effect of a myriad of cytokines and chemokines and adapt to the hypoxic niche, however fastidious continue to enjoy normal metabolism with no traces of Warburg effect in them[5].

There are many examples in which chronic inflammation is associated with carcinogenesis. However in none of them there is solid proof of a driver role for chronic inflammation[6,7]. In case of H. Pylori related maltona, early in the natural history of the disease cure can be obtained by eradicating H. Pylori with antibiotic[8]. Many people with chronic gastritis from H. Pylori never develop malignancy[9], demonstrating that chronic inflammation in and of itself is not sufficient to cause malignant transformation. Epstein Barr virus and its relation with Burkitt’s lymphoma in sub-Saharan Africa and nasopharyngeal carcinoma in China provide additional associations between inflammation and malignancy. Although inflammatory cells are observed in the tumor stroma, there is no proof of chronic inflammation as the driver which is best defined as an agent the removal of which would lead to resolution of the malignant phenotype[10].

Viruses can directly induce malignancy by down-regulating tumor suppressor genes such as TP53 and Rb. Through this mechanism papilloma virus promotes the pathogenesis of malignancies such as squamous cell carcinoma of cervix and head and neck[11]. The list of virus associated malignancies is quite long and includes...
Inflammation in hematologic malignancies

There is also evidence that chronic inflammation facilitates clonal evolution in myeloproliferative neoplasm (MPN), a chronic hematologic malignancy. Chronic inflammation is a characteristic feature of MPN and has been implicated as the cause of many of the debilitating symptoms associated with this disease. Emerging evidence suggests that inflammation may play a more central role in disease initiation and maintenance of the neoplastic clone16. Somatic acquisition of a JAK2V617F mutation in a hematopoietic stem cell is seen in the majority of patients with MPN. JAK2V617F confers upon hematopoietic progenitors resistance to the suppressive actions of TNF. An environment with high levels of inflammatory cytokines (such as TNF) selects for cells which have mutated in such a way to avoid these suppressive cues (such as the JAK2V617F neoplastic clone). The JAK2V617F mutant clone then expands and leads to unrestrained expansion of mature myeloid cells. The JAK2V617F neoplastic clone in MPN likely also induces an inflammatory state which further enhances its selective advantage. This inflammatory response elicited by the JAK2V617F clone may also be required for clinical manifestations of the disease. Clones with leukemia-associated mutations (such as JAK2V617F) can be detected in a few percent of normal aged individuals[17]. This suggests that presence of a clone with a leukemia-associated mutation is not sufficient to induce a clinical hematologic malignancy, and possibly inflammation is necessary to enhance the growth of the clone and lead to clinical disease.

Inflammation facilitates malignant transformation in glioma

There is evidence supporting inflammation as a tumor growth promoting but not initiating factor. One example is infiltration of glioma associated macrophage/microglia (MG). In CNS inflammation, MG have been shown to be capable of presenting antigens and activating T cells, thus playing a direct role in modulating brain inflammation[18]. A significant number of blood- born monocytes infiltrate into the tumor site[19]. Their trafficking to tumor sites has been proven to promote tumor growth by secreting mitogenic, angiogenic, and immunosuppressive cytokines rather than attacking glioma cells[20,21]. CCL2 (also known as MCP-1), a chemokine that attracts macrophages, is consistently overexpressed in a number of glioma cell lines and in some human high grade gliomas with MG infiltration[22,23]. PAX6 has been shown to play a tumor suppressor role in glioblastoma[24,25]. CCL2 expression in glioma cells was nearly abolished by overexpression of PAX6 (Figure 1). This demonstrates that in glioma inflammation facilitates malignant transformation.

Inflammation and free energy

One could arbitrarily categorize malignant disorders into two major groups. The first group would be those malignancies that are driven down an inflammatory path, such as the examples mentioned above. The second group would be those malignant disorders that are driven down the non-inflammatory path, such as certain well defined syndromes including Li-Fraumeni and Lynch syndrome, classical lung, breast and colon cancer among many others.

Free energy was first described by the American engineer Willard Gibbs in 1873 as the amount of energy available to do non-volume increasing work in a system with constant temperature and pressure and that inversely correlates with the amount of entropy of that system on thermodynamics arrow of time.

Biological systems including a living cell are well known for their unique capability to keep entropy at the lowest possible level dictated by the limits of the second law of thermodynamics. The breakdown of this fine balance which is the distinguishing hallmark of malignant disorders could be envisioned as the universal driving force capable of driving down both inflammatory and non-inflammatory roads of malignant transformation. The fine balance of the cellular energetics under normal condition is governing a broad spectrum of fundamental biological events extending from epigenome to Micro RNA network and incorporating dynamics of mitosis and homeostasis of cell cycle kinetics to DNA damage response and gene regulatory mechanisms[26,27]. Even though we might all be taken by surprise, chronic inflammation might simply be a reaction against low free energy cells and proteins and this could revolutionize our understanding of the pathogenesis of a broad spectrum of as yet puzzling disorders ranging from autoimmune and collagen vascular disorders to cancer. Clearly this represents a different perspective of the reality that we have become used to.

Interestingly enough even the contemporary literature has started to appreciate the interplay of metabolomics and inflammation[28]. Literature has not as yet looked at the potential role of the immune system in recognizing cells and proteins with lower free energy. Rather, the immune system has thus far been looked at as a system comprised of cells and humoral factors specialized in recognizing and destroying foreign cells and antigens as well as microorganisms. Even though HLA system and its biology have acted as the cornerstone of recognizing
self and non-self, the sheer discrimination between self and non-self has not been given any deeper and more fundamental attribute. This difference might simply be based on the difference between the free energy of these cells and proteins and self-cells and proteins. Most recently we have been witnessing a new wave of immune mediated strategies taking advantage of checkpoint inhibitors such as PD-1 and CTLA 4 antagonists in unleashing immune system against cancer[29]. Ex vivo training and modification of immune cells has also led to new treatment modalities such as provenge in metastatic prostate adenocarcinoma and CART in the treatment of acute leukemia[30,31]. Even though there is no cure in sight so far, the fact that unleashing the immune system could lead to eradication of malignant cells much more than the normal host cells which are totally different in their free energy supports this notion.

According to the universal free energy concept normal inflammatory cells with normal metabolism and energetics get attracted to the tumor sites comprised of a significant number of cells with significantly lower free energy/ lower in situ vibratory motion[2]. This attraction could be envisioned as an attempt at increasing regional free energy by destroying low free energy cancer cells. Many times cancer cells are equipped with the necessary means to disable the inflammatory cells. This inability of immune system leads to further augmentation of the inflammatory response inside the tumor microenvironment which is also responsible for some of the devastating constitutional symptoms that the cancer patient experiences. Indeed a significant number of solid malignancies station in regional lymph nodes and take advantage of the humoral survival and growth promoting factors inside the lymph node microenvironment before spreading hematogenously to other sites. However therapeutic recruitment of sophisticated immune cells to tumor sites which have significantly lower free energy could dramatically facilitate our future treatment strategies in treatment of cancer. For example if we could deliver nanotubes with the desired vibratory potential/

Figure 1: Suppressed expression of CCL2 (MCP-1) in the GBM cell line U251HF expressing transfected PAX6 (2.1, 2.2, and 2.3) as compared with non-transfected (p), vector transfected (vi, v6) and negative PAX6-transfected (2.6) cells. Cells were grown in DMEM/F12 under normoxic condition with serum deprivation for 24 and 48 hours before getting subjected to the assays. (A) Real-time qRT-PCR measurement of CCL2 expression normalized to ACTB and compared to the level in U251HF (p) with serum deprivation for 24 hours, using SYBR-Green master mix (Roche) and Light Cycler real time PCR instruments following methods described previously 1. (B) ELISA quantification of secreted MCP-1 from the same cells analyzed in A, using MCP-1 ELISA kit from Assay Designs (Michigan). Values include the mean + SD of 3 independent experiments. PAX6 level in each cell line / transfectants by western blot assay was shown in ref. 3.
jiggling loaded on immune cells which could dock on
cancer cells we might come with some breakthroughs
in our immune mediated cancer therapeutics. Rejection
of the transplanted organs could also be prevented
through similar refinements. Finally some of the famous
misunderstood drivers of malignant path such as RAS in
adenocarcinoma of pancreas, HER family in breast cancer,
EGFR in lung cancer and gliomas, all have one thing in
common, namely increasing the pace of mitosis of low
free energy cancer cells, a futile attempt at increasing the
free energy of cancer cells which could potentially lead to
inflammatory response in the tumor microenvironment and
promote debilitating cancer associated symptomatologies
such as fatigue and weight loss.

CONCLUSION

Although chronic inflammation has been noticed and
reported for a long time by many different observers both
in the vicinity of malignant cells and at the premalignant
lesions of a wide range of malignant disorders both of
hematological and solid subtypes, assigning a driver
role to chronic inflammation has faced insurmountable
technical as well as evidence based barriers.

CCL2 expression on the surface of glioma cells
which attracts macrophages to the tumor site, with its
ensuing activation of other components of immune
system through secretion of humoral factors on one end
and eradication of inflammation without any observable
effect on tumor growth and progression on the other end,
are among the examples that take away the designation
of driver role from chronic inflammation in inflammation
associated malignancies.

Available evidence supports a promoting role for
chronic inflammation following malignant transformation.
Interestingly enough at times the malignant cell itself is
the source of chemokines which attract the inflammatory
cell to tumor site as mentioned above. Finally in other
tumor types the malignant cells take advantage of the
nurturing microenvironment of the immune system organ,
namely lymph nodes before progressing into advanced
stages. Melanoma and breast carcinoma are among such
malignancies.

The modern generation of immunotherapy,
including CTLA4 and PD-1 antagonists, has enabled us to
make clinical achievements and prolong the life of patients
with metastatic melanoma.

However the toxicities associated with these
treatment modalities which originate in reacting against
the self-antigens at times have proven prohibitive. The
key question is if the immune system can recognize the
difference between the energetics of the tumor cell and
normal cell, why is it so incapable of destroying tumor
cells and in addition why does it contribute to promotion
and progression of malignant phenotype?

The answer to this question might reside in the
exceptional capability of the tumor cell to also change
the energetics of the infiltrating immune cells in favor of
a low free energy state which would disable these cells
as a result of modification of the quaternary structure of
its constituent functional proteins under low free energy
condition, in addition to taking advantage of their growth
promoting humoral factors. A new wave of improved
understanding and design of delicate measurement devices
of cellular energetics and the ability to reverse those
differences offers hope for future cancer therapeutics
modalities. Indeed even today we are witnessing a
 crude, however rewarding attempt at physical rather than
chemical cancer therapy by alternating electrical fields in
the treatment of recurrent Glioblastoma multiforme, called
NovoTTF-100A system which has been misunderstood
as only a change in electrical field of cancer cell without
appreciating the induced transient increase in free energy
of the cancer cell[32,33].

AUTHOR CONTRIBUTION STATEMENT

Dr. Fleischman wrote the paragraph entitled
Inflammation in hematologic malignancies.

Dr. Zhou wrote the paragraph entitled Inflammation
facilitates malignant transformation in glioma.

Dr. Afrasiabi created the concept and wrote the rest
of the article.

ACKNOWLEDGEMENT

We would like to acknowledge Mr. Matt Matsunari
from Twintel Solutions for assisting us in editing and
submitting the article at Genes & Cancer website.

CONFLICT OF INTEREST

The authors declare that they have no conflict of
interest whatsoever.

REFERENCES

1. Zhou, Y.H., Raj, V.R., Siegel, E. & Yu, L. Standardization
of Gene Expression Quantification by Absolute Real-Time
qRT-PCR System Using a Single Standard for Marker and
Reference Genes. Biomark Insights. 2010: 5: 79-85.

2. Afrasiabi, K.M.-S.,. J. Free energy measurement
distinguishes normal from cancer cell, offering a new
perspective for curing cancer. OnLine Journal of Biological
Sciences. 2013: 12: 11-125.

3. Zhou, Y.H., Wu, X., Tan, F., Shi, Y.X., Glass, T., Liu, T.J.,
Wathen, K., Hess, K.R., Gumin, J., Lang, F., Yung W.K.A.
PAX6 suppresses growth of human glioblastoma cells. J
Neuro-Oncology. 2005: 71: 223-229.

4. Warburg, O. On the origin of cancer cells. Science (New
York, N.Y.) 123, 309-314 (1956).
5. Kumar, S., Doni, T.R., Agnihotri, N. & Mehta, K. Transglutaminase 2 reprogramming of glucose metabolism in mammary epithelial cells via activation of inflammatory signaling pathways. International journal of cancer. Journal international du cancer. 2014: 134: 2798-2807.

6. Hasselbalch, H.C. Chronic inflammation as a promoter of mutagenesis in essential thrombocytopenia, polycythemia vera and myelofibrosis. A human inflammation model for cancer development? Leukemia research. 2013: 37: 214-220.

7. Saxena, M. & Yeretssian, G. NOD-Like Receptors: Master Regulators of Inflammation and Cancer. Frontiers in immunology. 2014: 5: 327.

8. Guo, Q., Guo, S. & Zhang, Y. Treatment of gastric MALT lymphoma with a focus on Helicobacter pylori eradication. International journal of hematology 97, 735-742 (2013).

9. Pilotto, A. & Franceschi, M. Helicobacter pylori infection in older people. World journal of gastroenterology. 2014: 20: 6364-6373.

10. Rickinson, A.B. Co-infections, inflammation and oncogenesis: future directions for EBV research. Seminars in cancer biology. 2014: 26: 99-115.

11. Friedman, J.M., Stavas, M.J. & Cmelak, A.J. Clinical and scientific impact of human papillomavirus on head and neck cancer. World journal of clinical oncology. 2014: 5: 781-791.

12. Mesri, E.A., Cesarsman, E. & Boshoff, C. Kaposi’s sarcoma and its associated herpesvirus. Nature Reviews Cancer. 2010: 10: 707-719.

13. Carbone, A., et al. Diagnosis and management of lymphomas and other cancers in HIV-infected patients. Nature reviews. Clinical oncology. 2014: 11: 223-238.

14. Zane, L. & Jeang, K.T. HTLV-1 and leukemogenesis: virus-cell interactions in the development of adult T-cell leukemia. Recent results in cancer research. Fortschritte der Krebsforschung. Progres dans les recherches sur le cancer. 2014: 93: 191-210.

15. Shermam, M. & Colombo, M. Hepatocellular carcinoma screening and diagnosis. Seminars in liver disease. 2014: 34: 389-397.

16. Fleischman AG, Aiehberger KJ, Luty SB, Bumm TG, Petersen CL, Doratotaj S, Vasudevan KB, LaTocha DH, Yang F, Press RD, Loriaux MM, Pahl HL, Silver RT, Agarwal A, O’Hare T, Druker BJ, Bagby GC & Deininger MW. TNFalpha facilitates clonal expansion of JAK2V617F positive cells in myeloproliferative neoplasms. Blood. 2011: 118: 6392-6398.

17. Xie, M., et al. Age-related mutations associated with clonal hematopoietic expansion and malignancies. Nature medicine. 2014: 20:1472-1478.

18. Hickey, W.F. & Kimura, H. Perivascular microglial cells of the CNS are bone marrow-derived and present antigen in vivo. Science. 1988: 239: 290-292.

19. Badie, B. & Schartner, J.M. Flow cytometric characterization of tumor-associated macrophages in experimental gliomas. Neurosurgery. 2000: 46: 957-961.

20. Chen X, Zhang L, Zhang Y, Liang J, Wang H, Ouyang M, Wu S, da Fonseca AC, Weng L, Yamamoto Y, Yamamoto H, Natarajan R, Badie B. RAGE Expression in Tumor-associated Macrophages Promotes Angiogenesis in Glioma. Cancer research. 2014: 74:7285-97.

21. Platten M, Kretz A, Naumann U, Aulwurm S, Egashira K, Isenmann S, Weller M. Monocyte chemoattractant protein-1 increases microglial infiltration and aggressiveness of gliomas. Annals of neurology. 2003: 54: 388-392.

22. Desbaillets I, Tada M, de Tribolet N, Diserens AC, Hamou MF, Van Meir EG. Human astrocytomas and glioblastomas express monocyte chemoattractant protein-1 (MCP-1) in vivo and in vitro. International journal of cancer. Journal international du cancer. 1994: 58: 240-247.

23. Goede, V., Brogelli, L., Ziche, M. & Augustin, H.G. Induction of inflammatory angiogenesis by monocyte chemoattractant protein-1. International journal of cancer. Journal international du cancer. 1999: 82: 765-770.

24. Zhou, Y.H., Hess, K.R., Liu, L., Linkskey, M.E. & Yung, W.K. Modeling prognosis for patients with malignant astrocytic gliomas: quantifying the expression of multiple genetic markers and clinical variables. Neuro-oncology. 2005: 7: 485-494.

25. Zhou, Y.H., Tan, F., Hess, K.R. & Yung, W.K. The expression of PAX6, PTEN, vascular endothelial growth factor, and epidermal growth factor receptor in gliomas: relationship to tumor grade and survival. Clinical cancer research. 2003: 9: 3369-3375.

26. Karr JR, Sanghvi JC, Macklin DN, Gutschow MV, Jacobs JM, Bolival B Jr, Assad-Garcia N, Glass JJ, Covert MW. A whole-cell computational model predicts phenotype from genotype. Cell. 2013: 150:389-401.

27. Jin, S., Li, Y., Pan, R. & Zou, X. Characterizing and controlling the inflammatory network during influenza A virus infection. Scientific reports. 2014: 4: 3799.

28. Kominsky, D.J., Campbell, E.L. & Colgan, S.P. Metabolic shifts in immunity and inflammation. Journal of immunology. 2010: 184: 4062-4068.

29. Smith, E.L., Zamarin, D. & Lesokhin, A.M. Harnessing the immune system for cancer therapy. Current opinion in oncology. 2014: 26: 600-607.

30. Singh, B.H. & Gulely, J.L. Therapeutic vaccines as a promising treatment modality against prostate cancer: rationale and recent advances. Therapeutic advances in vaccines. 2014: 2: 137-148.

31. Cheadle EJ, Gornall H, Baldan V, Hanson V, Hawkins RE, Gilham DE. CAR T cells: driving the road from the laboratory to the clinic. Immunological reviews. 2014: 257: 91-106.

32. Kirson, E.D., et al. Alternating electric fields arrest cell proliferation in animal tumor models and human brain tumors. Proceedings of the National Academy of Sciences.
of the United States of America. 2007: 104: 10152-10157.

33. Mrugala MM, Engelhard HH, Dinh Tran D, Kew Y, Cavaliere R, Villano JL, Annenelie Bota D, Rudnick J, Love Sumrall A, Zhu JJ, Butowski N. Clinical Practice Experience with NovoTTF-100A System for Glioblastoma: the Patient Registry Dataset (PRiDe). Seminars in Oncology: 2014: 41:S4-S13.