Cancer genes and cancer stem cells in tumorigenesis: Evolutionary deep homology and controversies

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Received 7 December 2021; received in revised form 10 February 2022; accepted 8 March 2022
Available online 4 April 2022

Abstract In the past, contradictory statements have been made about the age of cancer genes. While phylostratigraphic studies suggest that cancer genes emerged during the transitional period from unicellularians (UC) to early metazoans (EM), life cycle studies suggest that they arose earlier. This controversy could not be resolved. Phylostratigraphic methods use data from somatic tumor gene collections containing or lacking polyploidy genes (PGCC genes) and compare them to genes from evolutionary node taxa. I analyze whether the selected taxa are suitable to resolve the above contradiction or not. Both cancer and amoebae life cycles have a reproductive asexual germline that produces germline stem cells (GSCs) and somatic cell lines that cannot. When the germline loses its reproductive function, the soma-to-germ transition forms a new reproductive germline. The reproductive polyploidy of cancer is homologous to the reproductive polyploidy of unicellular cysts. PGCCs repair DNA defects, reorganize the involved genome architecture and produce new GSCs. The present study refutes the dogma of the early metazoan origin of cancer. Cancer has a unicellular life cycle that was adopted by early metazoans to rescue themselves from evolutionary dead ends. Early metazoans controlled the unicellular life cycle through suppressor and anti-suppressor genes that could suspend or reactivate it. They are the archetypes of tumor suppressor genes and oncogenes. Cells of mammalians and humans that reach a similar impasse as early metazoans can reactivate the conserved life cycle of unicellularians.

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Introduction

"Deep homology" is a recent concept in developmental biology and medicine and has its roots in evolution. Robert Kreisman noted in 2010 that all species are thought to have had the same ancestor hundreds of millions of years ago and that we all still share common, deep homologous genes. The same gene clusters (gene modules) that worked together millions of years ago still work together today. However, such gene modules are much easier to identify in less complex organisms than in human diseases or cancer. McGary et al studied more than 200,000 gene–phenotype associations in humans and lower animals and found matches in several species. Accordingly, scientists use a yeast model to study the angiogenesis of cancer and a worm model for breast cancer. All of these orthologous phenotypes reveal evolutionarily conserved gene networks. However, evolutionary models of human disease and cancer in their entirety are rare.

Over the last decade, I have studied the close similarity between the life cycle of Entamoeba and that of cancer. I found that both life cycles have many similarities and have summarized these in the "amoeba model" of cancer. Unfortunately, little is known about the ancestral gene module adopted by amoebozoans and metazoan cancers. However, there is no doubt that both cell systems share the same genetic machinery that goes back to the common unicellular ancestor. Not surprisingly, the life cycle of both systems consists of a reproductive germline and a somatic cell line (G+S life cycle). This is due to the fact that amoebozoans (including Entamoeba) and Metazoa (including cancer) are the last branched sister clades of the phylogenetic tree of life. Both belong to the group Amorpha.

I disagree with the common dogma of the origin of cancer by early metazoans and argue that cancer is an expression of a pre-existing life cycle that was adopted by unicellular organisms, but also by early metazoans from their common ancestor to escape from evolutionary dead ends. I assume that early metazoans already controlled the adopted unicellular life cycle by suppressor and anti-suppressor genes and that these are the archetypes of tumor suppressors and oncogenes.

After branching, the unicellular G+S life cycles of Amoebozoa and early Metazoa cycles evolved separately. They optimized their genetic machinery in response to the environment. Entamoeba and metazoan cancers evolved to the parasitic lifestyle that optimally exploited host resources and niches. However, both systems retained the basic structure of the ancestral cell system and passed it on to subsequent generations without fundamental changing. This explains the deep evolutionary homology in terms of germline reproductive polyploidy, stemness, stem cell phenotypes, and stem cell repair, and suggests a more coherent evolutionary designation for participants in the life cycle of cancer.

Using phylostratigraphic methods and analyzing data from somatic tumor gene collections that contain or do not contain polyploidy genes from PGCCs and taxa with evolutionary nodes, the present work shows that cancer polyploidy is homologous to the deep polyploidy of the unicellular ancestors that evolved stemness and the first archeal stem cells. The mechanisms of stem cell formation by reproductive polyploidization were adopted by protists such as Entamoeba as well as metazoan cancers.

Little is known about the evolutionary origin of cancer stem cell (CSC) and their archetypes, so much of this work is concerned with the evolutionary origin of CSCs. I argue that one of the crucial events in the evolution of pre-metazoans was the development of the inner cyst cell into a stem cell progenitor phenotype and founder stemness genes. The evolution of CSC archetypes traces back to the common ancestor that transformed the simple protective cyst of the ancestral flagellate into a reproductive cyst capable of stemness. Its haploid daughter cells transferred stem cell potential into the amoeboid stem cells that differentiate the germline. The germline of the common ancestor evolved to carry stemness.

This review begins with a brief introduction to the evolutionary "deep homology" and considers cancer evolution from the perspective of deep homology. It explains the cancer life cycle using the "amoeba model" and shows that CSCs are products of their unicellular life cycle of cancer. It concludes that the origin of the ancestral inherited life cycle is found in the polyploid reproductive cysts of single-celled organisms.

It can be assumed that cancer gene archetypes evolved in the pre-metazoan era in the form of a Gaussian bell diagram. Many G+S founder genes are cancer gene archetypes that arose early in evolution when simple protective cysts evolved into reproductive cysts that became polyploid. Last but not least, this work addresses the question of why metazoans did not completely abandon the unicellular G+S cycle. I suggest that unstable early metazoans that carried the G+S cycle for safety reasons suppressed it only to the extent that reactivation remained possible when evolutionary dead ends arose. This was beneficial for the whole-cell community as well as for individual cells. In the early era of metazoans (EM), new EM suppressor genes

| Abbreviations |
|---------------|
| UC | unicellularians |
| EM | early metazoans |
| MC | multicellularians |
| MM | mammalian |
| MGRS | multinucleate genome repair structures (cancer); multinucleate genome repair syncytia (Entamoeba) |
| PGCC | polyplod giant cancer cell |
| aCLS | autonomous cyst-like structure |
| GSCs | germline stem cells |
| G+S | germ and soma |
| GST | germ to soma transition |
| SGT | soma to germ transition |
| EMT | epithelial—mesenchymal transition |
| MET | mesenchymal—epithelial transition |
| GRN | gene regulatory network |
| DDR | DNA damaged repair |
| SSB | single-strand break |
| DSB | double-strand break |
| EAS | early surface antigens |
were integrated into the EM regulatory network, but also UC genes that were repurposed as anti-suppressor EM genes. This regulatory hub gene module allowed metazoans to control the G+S cycle and retain it as useful.

Deep homology, the ancestral gene module and cancer evolution

According to Shubin et al, "deep homology" means an ancient similarity of patterning mechanisms and a historical continuity through which regulatory circuits inherited from a common ancestor have been passed on in evolution. This means that new functions and phenotypes have not emerged from scratch; they have evolved by exploiting regulatory circuits that were already established in early eukaryotes. As Wagner added, individual structures that do not appear to be homologous at first consideration may have evolved from the genetic machinery of a common ancestor, using it as a pattern for homologous evolutionary programs. In recent years, deep homology approaches have evolved more and more towards a better understanding of the molecular and genetic mechanisms and function, moving away from the simple comparative morphology.

Tschopp and Tabin understand deep homology as the conservation of gene expression during evolution and speak from historical continuity, historical homology, and organizational hierarchy. They consider the homologies between the individual cell types of a given organ as the core of the modern concept of homology and assume that phenotypes result from complex gene regulatory networks (GRN) and gene modules rather than from single individual genes. According to the "kernel concept" of Davidson and Erwin regarding the hierarchy in the regulation of genes, kernels are sub-units of GRN that exhibit deep evolutionary preservation and are refractory to regulatory rewiring and underlie stability while at the base of the GRN hierarchy are so-called gene differentiation batteries that direct terminal cell differentiation.

The evolution of clade genomes began with the evolutionary branching 750–800 million years ago and continues until today. Over time, ancestral genes have taken on additional or entirely new functions, adapted to analogous microenvironmental pressures in the tissue they parasitize, and passed these changes on to subsequent generations of cells, according to the Darwinian model of evolution. Recent work by Vendramin et al shows that the Darwinian model alone is insufficient to fully explain cancer evolution, as "single catastrophic events such as whole-genome duplication (WGD) position cancer evolution beyond Darwin". It confirms Shubin's assumption that innovations have a history that goes far back deep in time. However, I think, WGD is not a catastrophic event, rather, it is a reproductive process inherited from the common ancestor.
Cancer evolution through germline clones and subclones is certainly beyond Darwin. It is largely related to deregulated processes of soma-to-germ transition (SGT, EMT) caused by various environmental stress factors. Environmental stress leads to altered and incomplete second CSCs generations and multiclonal germline stem cells (GSCs). Cancer evolution is in contrast to the previously assumed linear sequential evolution of clonal cell division. Gradual changes in CSC progenitors provide cancer with a strong selective advantage, allowing earlier CSCs lineages to be displaced.14–17

The life cycle of cancer explained by the “amoeba model”

The great similarity between the unicellular life cycles of Entamoeba and cancer contradicts the molecular phylogenetic theory for the early metazoan origin of cancer.18–21 The deep relationship between the two cell systems, resulting from the temporally close branching of Amoebozoa and Metazoa (Amorpha group), is why the “amoeba model” is a uniquely valuable support for understanding the biology of cancer from the evolutionary perspective.22

The G + S life cycle of Entamoeba is closer to the common ancestor and more original than any other life cycle of unicellularians. Both cell systems — amoeba and cancer — use the deep homologous G + S gene module evolved by the common ancestors. Both systems have striking homologies: (i) A reproductive asexual germline capable of forming germline stem cells (GSCs, referred to as CSCs in cancer) and a somatic cell line without reproductive GSC function; (ii) Germ and soma cells that proliferate through asymmetric and symmetric cell cycles and can interconvert by transitioning from germ to soma (GST) and from soma to germ (SGT); both processes are referred to as MET and EMT in cancer; (iii) Oxygen-sensitive germlines that irreversibly lose their reproductive function due to irreparable DNA damage caused by excess oxygen; (iv) DNA damage repair (DDR) mechanisms to repair DNA replication and polyploidization defects and maintain genomic integrity of nascent GSCs/CSCs; (v) DNA DSB repair mechanisms via MGRS and PGCC structures, with or without homologous cell fusion; It is also conceivable that the release of excess DNA material, as occurs in Entamoeba by MGRS or non-reproductive polyploidization/polyploidization processes, results in circulating cancer DNA fragments.

In Entamoeba, asexual reproductive cycles begin with the polyploidization of the inner cyst cell, which gives rise to eight haploid daughter cells that develop into diploid stem cells (Fig. 2). I refer to these “nascent” stem cells produced by the germline as GSCs. These GSCs differentiate into the reproductive germline that proliferates by asymmetric cell division. This type of cell division gives rise to non-identical daughter cells: one of them is the self-renewing SR daughter cell (D1 cell) and the other the non-proliferating sister cell (D2 cell) committed for reproductive polyploidization and GSC formation. Each D1/5R cell continues the normal cell cycle while D2 cells leave it to transform into a reproductive cyst. The inner cyst cell changes its structure and undergoes several whole cell cycle rounds (WGDs) without cytokinesis and cell division. The closed asexual circuit of Entamoeba’s germline includes cysts, GSCs, self-renewing germline cells (D1), and cyst precursor cells (D2). In cancer, the term GSCs used for nascent CSCs has the advantage of clearly indicating the origin of primary CSCs.

Cancer stem cells are products of the G + S life cycle

Cancer stem cells are products of the G + S life cycle and are not derived from adult HSCs or de-differentiated cells as previously thought. They are products of the unicellular G + S life cycle adopted from early metazoans. Cancer is ubiquitous in metazoans,6 including low metazoans such as Holozoa and other lower animals. For 700–800 million years, metazoan cancers have had reproductive germlines that performed polyploid aCLS cycles via encystment (reproductive germline cycles) and produced GSCs. Nascent CSCs are GSCs of unicellular origin and, like Entamoeba GSCs, are defined by the asymmetric cell cycle, asymmetric cell division, and differentiation ability. During the evolution of cancer, the germline was involved in the formation of different classes of CSCs which are: (i) the nascent GSCs (primary CSCs), (ii) the SGT/EMT-induced CSCs (secondary CSCs), and (iii) the MGRS/PGCC-induced CSCs (tertiary, repair CSCs). They correspond to the GSC classes and mechanisms of GSC production observed in Entamoeba.

In cancer, the reproductive germline cycle begins with the committed D2 cell (precursor cell) that polyploidizes within a cell envelope thinner than the cyst wall. The cancer germline undergoes development similar to the Entamoeba’s germline. There is a deep homology to the mammalian GSCs. The amoeba model substantiates the statement of Nayernia, who hypothesizes that mammalian (somatic) stem cells are the direct descendants of the germline.23,24 According to this hypothesis, the germline is the “mother line” of all somatic stem cell lineages in the adult body and the daughter GSCs are the only stem cells capable of passing genetic information from generation to generation.23,25

The germline of amoebae and cancer is oxygen-sensitive. Under conditions of excess-oxygen it loses its reproductive function and cannot produce nascent CSCs/GSCs (Fig. 3). As a result, soma cells are forced to form new reproductive germline clones that begin with the production of secondary sCSC/sGSC.4 When soma-to-germ transition (SGT/EMT) cannot be initiated, defective germline cells repair DNA defects and reorganize the genome architecture of the germline through multinucleate genome repair structures (MGRSs) or polyploid giant cancer cells (PGCCs) with or without cell fusion.

In the amoeba model, MGRS/PGCC polyploidization does not activate ancient developmental programs, as previously claimed.26,27 This occurs during the transition from the weakened multicellular life cycle of metazoans to the unicellular G + S life cycle of cancer. During cancer development, polyploidization occurs not only in MGRS/PGCC structures induced by chemotherapeutic agents or irradiation, but also in the early stages of cancer, when excess oxygen damages the germline. The damaged
germline cells have lost their reproductive function and cannot form new CSCs. To repair DNA defects and remodel genome architecture, oxygen-damaged germline cells fused and formed autonomous MGRS/PGCC syncytia that repair DNA DSB damage and produce new effective CSCs capable to differentiating new reproductive germline clones. I disagree with the notion that polyploidy in cancer is an important driver of epigenetic change and that "atavistic" changes go hand in hand with cancer polyploidy. Rather, the "amoeba model" confirms the assumption that cancer "defects" (such as reproductive polyploidization) are mostly adaptive tools of survival programs that evolve during the evolution of life and are inherent to cancer.

The deep homologous origin of the $G+S$ life cycle of cancer lies in the reproductive cyst of unicellularians

Crucial to the evolution of unicellularians (UCs) was the transition from simple protective cysts to reproductive cysts capable of polyploidy. Cyclic polyploidy is the result of the evolution to asymmetric cell division. According to Schussnig, the archetype of the last eukaryotic common ancestor LECA was a mononuclear and motile "monad cell" that could form cysts as a defense against harmful living conditions. Within this protective envelope, the flagellar cell reduced its morphology to an amoeboid cell shape, which left the cyst wall after a resting phase. This inner cyst transformation occurred with successive regression of the flagellar phenotype. While early eukaryotic cysts were non-reproductive structures, later eukaryotic cysts did not cease replication and performed multiple rounds of WGDs without cytokinesis and cell division. Such polyploid cysts gave rise to more than two daughter cells. Polyploidization, initially random and facultative, became systemic due to better progeny survival and an evolutionary advantage.

The transition from protective to reproductive cysts had great evolutionary potential. In addition to phenotype transition, the reproductive function was transferred from the flagellar to the amoeboid cell type and its daughter cells. These germ cells were generated by polyploid cycles within the reproductive cyst, and the flagellar cells express only the somatic genome. The best evidence for this is provided by Proleptomonas, a free-living, mononuclear flagellate with a reproductive cyst $6-9 \mu m$ in diameter that belongs to the Cercozoa. Unlike other flagellates that divide vegetatively by symmetrical cell division, Proleptomonas reproduce only within the reproductive cyst and, like Entamoeba, produces eight daughter germ cells that form the germline. Early amoeboflagellate ancestors have laid the groundwork for germ and soma life cycles. The flagellate cell transferred its totipotency to the innercyst cell and its amoeboid daughter cells, which gradually took on stemness and differentiation capabilities, in contrast to the flagellate phenotype, which expressed only soma traits. Both phenotypes retained the capacity for inter-conversion and could transit from germ-to-soma (GST) and from soma-to-germ cells (SGT). The amoeboflagellate ancestor transferred the deep homologous $G+S$ gene module to the branching taxa (Fig. 1) that adopted either the entire gene module, or only parts of it in active or inactive forms, or discarded them altogether. With rhizopodial transformation, pre-metazoans expanded the possibilities for colonial life, cell fusion, and lateral gene transfer (LGT). LGTs enable the asexual exchange of genetic information between different genomes.
Accordingly, it can be assumed that the common ancestor of *Entamoeba*, Metazoa and cancer was a highly evolved amoeboflagellate with a relevant deep G + S module including genes for stemness and cell differentiation, polyploidy as well as GST and SGT processes. Sponges (Porifera), which belong to the oldest metazoans, support this hypothesis. Stem cells and cell differentiation in modern sponges arise from totipotent amebocytes (archeocytes) (Fig. 4) and not from flagellated choanocytes, which have only soma function and differentiate themselves from amebocytes. Nevertheless, SGT processes are also known in sponges, where soma choanocytes transform into totipotent amebocytes. 31

**Phylostratigraphic errors: choanoflagellates are not the taxa closest to metazoans**

Conventional wisdom holds that choanoflagellates are the unicellularians that would be closest to metazoans. But this seems to be less and less true. Choanoflagellates and Amoebozoa are two groups to Metazoa. As revealed by Carr et al, little is known about choanoflagellate evolution. 32 Unlike Amoebozoans, Choanoflagellates show a low propensity to form cysts and dual life cycles and there is no evidence that choanoflagellates have an exclusive relationship with Metazoa. 33 Molecular data reject suggestions that Metazoa are descended from a true choanoflagellate ancestor. Much more is conceivable that both groups are descended from a common protistan ancestor. From this ancestor, metazoans evolved into truly multicellular organisms, whereas the choanoflagellates have maintained a predominantly solitary existence.

Recent molecular studies showed that the transcriptome of choanocytes bears little resemblance to the transcriptome of choanoflagellates. 31 This finding clearly refutes the widely held view that metazoans evolved from a unicellular ancestor surrounded by an apical cilium, and a microvillar collar structurally similar to modern choanoflagellates. 34–36 Brunet and King hypothesized that the first metazoans evolved from a colonial state of unicellular organisms with a long history of "simple multicellularity" that captured nutrients and bacteria with a flagellum surrounded by a microvillar collar and had a...
Figure 4 Sponges architecture: choanocytes (red) and amoebocytes/archeocytes (green). From: Ruppert EE, Fox RS, Barnes RD. Invertebrate Zoology, ed. Brooks/Cole. p. 82. ISBN 978-0-03-025982-1 (https://en.wikipedia.org/wiki/Sponge).

The dogma of the early metazoan origin of cancer

In 2007, Domazet-Loso and Taz introduced phylostratigraphic analysis to study human genome history and the evolutionary origin of genes related to human genome diseases, including cancer.\(^\text{18,19}\) Central to phylostratigraphic research is the assumption that new protein families that emerge at evolutionary nodes — such as the transition to multicellularity — are the major innovations for life initiated by founder genes. The aim of the phylostratigraphic analyses was to find out whether there is a link between the founder genes of multicellularity and cancer genes.

The phylostratigraphic map of human genes of Domazet-Loso and Taz has 19 phylostra (ps)\(^\text{19}\) (Fig. 5). The first three layers harbor archaea and bacteria (ps1), lower eukaryotes (ps2), and fungi (ps3), while on the fourth phylostratum researchers placed the less adequate choanoflagellates, a decision that led to misinterpretation. Moreover, the analyzed gene material was derived from somatic cell material of tumors and metastases and was accordingly poor in CSC gene material, which could have provided more information on the age of the cancer's germline genome. Tumors and metastases are poorly populated with CSCs. The number of functional CSCs can decrease progressively and rapidly in tumors.\(^\text{12}\) After about 100 cell doublings and a spontaneous extinction probability of 95%, the metastatic efficiency decreases to about 0.05% (loss of function).

Based on this map, the researchers found a significant number of cancer protein domains and cancer-like genes mainly at the level of ps1 (Archaea and Eukarya), followed by a decrease on ps2 and ps3 (primitive Eukarya and Fungi) and a complete missing in choanoflagellates (ps4). A second significant peak occurs in the early metazoans of ps5 (Porifera) (Fig. 5). The occurrence of cancer-like genes was significantly reduced at the ps6- and ps7-levels (Eumetazoa and Bilateria) and at all levels beyond the emergence of vertebrates (ps11 and up).\(^\text{18,19}\)

The thinking at the time was influenced by Weinberg, who assumed that cancer is a consequence of multicellular evolution.\(^\text{42}\) However, the peak observed in ps1 predates the emergence of multicellularity. Domazet-Loso and Taz believe that the deep peak indicates primitive protein domains associated with so-called cancer "caretaker" genes that arose during the formation of the first cells to ensure the stability of the genome. In contrast, the overexpressed ps5-peak would be due to the so-called "gatekeeper" genes, and contains oncogenes- and tumor suppressor-like
Figure 5  Statistical analysis of the cancer datasets using the 19-phylostrata map. From: Domazet-Lošo and Tautz \cite{DomazetLošo2018} (http://www.biomedcentral.com/1741-7007/8/66). CC BY 3.0.
genes. In the ontogenetic sequence of tumor progression, mutations in caretakers precede mutations in gatekeepers.

In more recent phylostratigraphic studies, Trigos et al reduced the number of phylostrata to 16 and grouped all human genes into three super-classes: 6719 unicellular ones (UC) originating from the phylostrata ps1–ps3, 7939 arising from the early metazoan (EM) of the phylostrata ps4–ps9, and 2660 genes arising from the mammalian phylostrata ps10–ps16 (MM). Accordingly, ancient genes related to cancer (ps1–ps3) are termed UC genes; they include genes of Opisthokonta (flagellated protozoa) and choanoflagellates, but not amoebozoans. The authors suggest that UC genes support basic cellular functions such as cell cycle, mitosis, proliferation and metabolic processes. In contrast, genes related to cancer (ps4–ps9), are referred to as EM genes; they represent genes with more complex cellular functions that arose with the transition to multicellularity.

**Phylostratigraphic polyploidy studies reject the dogma of the EM origin of cancer**

In an attempt to determine the evolutionary age of cancer polyploidy genes, the research group around Erenpreisa and Anatskaya analyzed the occurrence of polyploidy genes in the UC, EM, and MM phylostrata. For this study, the researchers used transcriptomic data from tumors and tissue containing polyploid PGCCs. These PGCCs are multicellular genome repair structures (MGRSs) capable of repairing defective germline cells and reorganizing the germline genome architecture. PGCCs generate new invasive CSCs capable of metastases.

The results in Figure 6 show that genes related to cancer polyploidy arose quite early in the evolutionary history. Both ploidy-related and bivalent ploidy-regulated genes (BVGs) belong to ps2 and occur at a time when primitive eukaryotes have switched from simple protective cysts to reproductive polyploid cysts. In contrast to the previous results of Domazet-Loso and Taz, the distribution of BVG genes show the expected Gaussian bell curve, ranging from ps1 to ps7. It is unfortunately interrupted by the miss-selected choanoflagellates and opistochonts on ps3. The curve flattens with the appearance of metazoans and finds its lowest point in the phylostrata representing Bilateria and Chordata (ps6/ps7).

It is clear that the work of Anatskaya et al traces cancer-related polyploidy genes to a time when the unicellular ancestor was developing reproductive polyploid cysts, and paving the way to stemness, cell differentiation, and G + S life cycles, overturning the dogma of the EM origin of cancer. The relationship between the reproductive polyploidy of cancer and the reproductive ploidy of ancient cysts is a clear supportive argument for cancer’s unicellular origin. However, I disagree with the idea of the authors that polyploidy could “activate the recapitulation of evolutionary developmental programs.” In my opinion, cancer polyploidy is firstly (i) a reproductive mechanism of intact germlines and clones to generate nascent CSCs (cyclic reproductive polyploidy) and secondly (ii) a repair and regeneration mechanism by which therapy-induced PGCCs renew damaged germline cells and CSCs pools. In both cases, polyploidization takes place within the G + S life cycle. Polyploidization is not the driver that switch human cells into the unicellular G + S lifestyle; it is part of the already reactivated unicellular G + S life cycle and a distinct feature of unicellular lifestyle.

**Suppression mechanisms of G + S life cycles evolved in early metazoans**

Multicellularity could not prevail unless the unicellular G + S life cycle could be prevented. Mechanisms blocking the unicellular life cycle were required to maintain multicellularity. In this sense, early metazoans evolved gene regulatory networks (GRNs) that consisted partly of newly evolved EM genes, but in large part the UC genes were repurposed as anti-suppressor genes (oncogene archetypes). The goal of the antisuppressor genes was to shut down the suppressor networks (tumor suppressor archetypes) and restore the unicellular G + S life cycle. The transition period from UC to EM was the era of constant alternation between unstable EM states and reactivated UC states.

In tumors, there is much molecular evidence of ancient UC and EM genes or gene modules that are upregulated or downregulated during tumor development. Phylostratigraphic research has shown that the UC subnetwork of the human genome is connected to the multicellular subnetwork (MC) through “hub” genes that, in healthy
Evolutionary deep homology and controversies

1243

cells, control and regulate the equilibrium between UC and MC genes. 21,27,43 This hub gene network corresponds to the suppressors and anti-suppressor genes emerging in the EM era. According to phylostratigraphic analyses, anti-suppressor genes, in metazoans and humans, disrupt the principles of UC/MC cooperation, especially in advanced neoplastic stages and metastases. They inactivate the MC genes, and upregulate the UC genes. 23

Genes of UC and MC origin are involved in numerous processes of healthy human cells. In the somatic tumor cells molecular researchers identified a set of deeply homologous genes as regulators of the coexpression of the UC and MC subnetworks. 21 Among this hub gene network, researchers found three UC genes from the phylostratum ps1 (ACTG1, RCC2, PKN2), five UC genes from ps3 and ps4 (PLEC, TLN1, VASP, DSP, CTNN ), and four EM genes from ps5 (ILK, CTNN1A1, CTNND1, and PKP3). Because of their central role in the human gene network, these 12 genes have fundamental vulnerability and play critical roles in cancer processes associated with genomic instability, late tumorigenesis, and metastasis. 21,43 However, it is not sure that hub genes play a crucial role in cancer initiation and the conversion from a normal, healthy cell into malignant cells as claimed by previous authors. 44–46

In tumors, Trigos et al found 12,812 downstream target genes and 1370 regulator genes and hypothesize that regulator genes control a variety of target genes. 21 Over half (56.42%) of the GRN genes are EM genes, while only 37.88% and 5.69% are UC and MM genes, respectively. The authors looked for genes with key regulatory roles controlling at least 10 downstream targets and designated them as master regulators. They found that 65.12% of the master regulators are EM genes, while only 28.49% and 6.40% are UC and MM genes respectively, and consider that the most important master regulators in the network are EM genes.

Genetic and non-genetic CSC heterogeneity arises from deregulated EMT processes

Posttherapeutic changes in the tumor microenvironment such as cell senescence and other environmental changes can alter both the phenotype and function of CSCs. However, CSCs may also lose their function prior to treatments if exposed to unfavorable growth factors such as hyperoxia (Fig. 3). Hypoxia-grown germlines suffer DNA DSB defects. They signal their loss of function to soma cells, which respond by soma-to-germ transition (SGT/EMT) and generate new functional germline clones. This is a process of germline renewal inherited from unicellular ancestors. The “amoeba model” provides an evolutionary explanation for cancer cell theories such as “cellular plasticity” and “clonal evolution”, which state that nonstem cells and stem cells can interconvert and that genomic and genetic alteration accumulate over time, leading to more CSC heterogeneity and increased aggressiveness. 47–50

In the amoeba model, somatic cells arise (i) either from stem cell differentiation or (ii) from germ-to-soma transition (GST). Somatic cells have epigenetically silenced the germline genome but can reactivate it when the germline signals DNA damage and loss of reproductive function. SGT is also one of the causes of increased pathogenicity in parasitic Entamoeba. In cancer, germline signaling induces SGT/EMT that forms secondary germlines and clones. EMT-generated CSCs are usually more pathogenic, leading to metastases.

SGT/EMT processes experience both epigenetic and genetic alteration. As a result, genetically distinct CSC clones coexist in different tumors. 51–55 They are differentially radiation sensitive and exhibit increased resistance to cancer therapies. Therapeutics can lead to adaptive responses and acquired radiation resistance and even more invasive tumors. Therapeutics preferentially kill non-stem cells and enrich CSC density in tumors from less than 1% to more than 80%, 51,56,57 CSC clones with genomic alteration protect tumors from therapeutic effects. Second, the ability to repair DNA damage after irradiation, particularly severe DNA DSB damage, is higher in CSCs than in their nontumor counterparts. 58 CSCs preferentially repairs DNA damage through ancient mechanisms of homologous recombination (HR) and MGRS/PGCC pathway. 59,60

Acquired resistance to anti-cancer therapy usually results from nongenetic alteration such as DNA methylation, histone modifications, and other epigenetic mechanisms that alter gene expression without changing the genetic profile. Such mechanisms act more rapidly than genetic mutations. 61–63 Although patterns of DNA methylation are generally faithfully transmitted to daughter cells, they can also be erased resulting in a large reduction in total DNA methylation. 64

Cancer stem cells generated by germline have three distinct superclasses, for which I have proposed the designations (i) nascent CSCs or primary GSCs; they arise from repetitive aCLS cycling and cyclic reproductive polyploidization; (ii) EMT-generated CSCs (secondary CSCs) that arise when the damaged germline signals the loss of reproductive function to soma cells, and (iii) PGCC-generated CSCs (tertiary CSCs), arising from repair and genome reorganization processes that occur in both native PGCCs (MGRS) and therapy-induced PGCCs. 64–66 (Fig. 3). PGCCs have mechanisms of DNA restructuring and natural genetic engineering capacities necessary to rearrange genomic components and its system architecture. All CSCs are produced from the germline. Late EMT enables tumors to become more malignant and metastatic. Initiation, maintenance, and inhibition of EMT are regulated by small molecules of antiproliferative agents. 65 External signaling molecules and intracellular signaling molecules activate various EMT transcription factors and mesenchymal EMT markers.

EMT processes involve the transition from specific epithelial markers (soma cell markers) to mesenchymal markers (CSC markers). Several transcription factors and other biomarkers are also involved in the orchestration of EMT that includes concomitant deletion, upregulation, and gene overexpression. Numerous cancer researchers agree that CSC formation is associated with EMT activation and that EMT leads to stem-like features and phenotypes that promote tumorigenesis and accelerated metastasis. 66–68

Some of these CSCs express both soma and germline markers simultaneously. They are considered to be alternative CSC-states or CSC-like states, conditioned by partial EMT processes. Such intermediate EMT stages are better known as hybrid, early hybrid, and late hybrid EMT stages.
The EMT-related CSCs are commonly referred to as "stem-like phenotype," "stem- and EMT-based gene expression," or "stem-like state" leading sometimes to confusion. Regardless of their origin, these are all cells with stemness function that have emerged from an earlier germline or a later germline with more or less invasive and metastatic potential.

In recent years, EMT has been increasingly understood as a common regulator of CSC phenotype, CSC production, and drug resistance. All in all, EMT is a critical mechanism leading to more malignancy and metastasis. It allows flexible gene expression between activation and repression. In contrast to the SGT processes in Entamoeba, cancer EMT are considerably deregulated by the immuno-suppressive tumor microenvironment.

Conclusions and perspectives

In recent years, the prevailing somatic mutation theory (SMT) of tumorigenesis has come under increasing pressure and several alternatives have been proposed and three main theories of cancer stem cell origin and CSC development have gained popularity. Each of them targets different phases of CSC evolution. One is the somewhat older "CSC model" theory, which proposes a unidirectional development model. This theory states, among other viewpoints, that primary CSCs divide asymmetrically to form a CSC pool and switch to symmetric cell cycling to increase the number of CSCs. The "clonal evolution model" assumes an accumulation of genomic and genetic alterations over time, leading to an increase in CSC heterogeneity and aggressiveness, while the "cellular plasticity model" assumes that non-CSCs and CSCs can interconvert. These theories are not mutually exclusive, but they address only part of the question. In contrast, the "amoeba model" of cancer, which I propose in the present paper, considers the evolution of cancer stem cells from the perspective of the "deep homology" of G + S gene module and its transition from a free-living cell system to a parasitic lifestyle which also occurs in Entamoeba. The "amoeba—cancer model" combines all three theories mentioned above into a more comprehensive theory of evolution. It distinguishes between "healthy" germline cells, which proliferate through asymmetric cell cycles and express stemness potential, and damaged germline cells, which lose their stem potential (loss of function) and proliferate through symmetric cell cycles.

Cancer ancestors were unicellularians evolving over millions and millions of years into sophisticated G + S life cycles. The origin of the deep homologous G + S gene module is in Ur-flagellates that could transform the protective cysts into reproductive cysts. The progeny of reproductive cysts evolved into stem cells, which differentiate germ and soma cell lines and cells that are capable of phenotype conversion. Damaged germline cells that lost the reproductive potential have learned to repair DNA defects and reorganize the genome architecture through MGRSs and/or PGCCs rearrangements. As colonial unicellular organisms or more advanced early metazoans began to evolve the integrated multicellular mode of life, evolution retained the old unicellular G + S life cycle as a stable protective measure for the maintenance of life. The extension of the GRN by new EM genes and repurposed UC genes took over the control and surveillance function over the ancient G + S gene module and its reactivation in case of evolutionary dead-ends. The extended GRN emerged suppressor and antisuppressor genes as archetypes of tumor suppressor genes (TSGs) and oncogenes. In parallel with the evolution of metazoans, the G + S gene module evolved the more complex cancer G + S life cycle contained in the human genome. From an evolutionary point of view, the G + S life cycle of unicellularians was not abandoned by metazoans, it was maintained as an "ultima ratio" in the genome of all metazoan and human cells.

The present work is one more analysis of the close relationship between the germline of cancer and the nascent cancer stem cells (nCSCs). I distinguish between three classes of CSCs: (i) nascent CSCs or primary CSCs are germline stem cells (GSCs) that arise from reproductive germine cycles; (ii) EMT-induced CSCs and EMT-induced CSC-like cells arise from somatic cells that transit into new germine cells, and clones through SGT/EMT processes; and (iii) high metastatic MGRS/PGCC-induced CSCs. These tertiary CSCs are the result of DNA repair and genome reorganization processes. Some SGT/EMT processes may be incomplete, defective, or deregulated, resulting in a broad spectrum of heterogenous CSCs and CSC-like cells that are either pure germline cells or hybrid cells with mixed germline and soma hallmarks.

Non-genetic sources of phenotypic variation may play an important role in tumor initiation, tumor progression, and evolution.

Recent research does not provide a definitive answer to the question of what cancer is. Is it an "atavism," a "genetic disease," or simply an ancient life cycle adopted by early metazoans to avoid evolutionary dead ends? It is now clear that cancer is not an atavism, as has been erroneously claimed in the literature in recent years. According to the definition by Wagner, an atavism is a morphophysiological trait controlled by an ancient gene regulatory network (GRN) that survives periods of disuse and can be reactivated and reused later lineages, even if it was not used in the immediate ancestors. Cancer is not an atavism because it was reactivated and reused continuously in the last 800 million years (Ma) by all intermediate ancestors of humans, mammalians, and vertebrates.

To summarize: What we perceive as an extremely unpleasant and complicated human disease (cancer) is a rescue mechanism of a sick, weakened metazoan cell that cannot continue its multicellular life and find itself in the same dead-end situation as their early metazoan ancestors 800 million years ago. This rescue mechanism, which reactivates the framework of the deep unicellular G + S life cycle, leads to a kind of aggressive pseudo-parasitism within the multicellular host organism and eventually to its death, which parasitic Entamoeba also do. However, Entamoeba eliminates stem cell precursors via cyst excretion, whereas cancer, as an ancient cell system closed in the host organism, maintains all nascent CSC generations that further potentiate the disease. In a previous paper, I proposed that severely threatened cells are pulling old DNA repair mechanisms out of Pandora's tool box for unicellular organisms, inadvertently restarting the dormant G + S life
cycle doing more harm than good. In many countries around the world, nearly one in two people will develop cancer once in their lifetime. A large variety of metazoans are also affected by cancer.

Correctly understanding the origins and mechanisms of using the "amoeba model" can speed progress towards preventing or curing cancer. Interestingly, there is older evidence that Entamoeba cells change their surface antigen architecture when transferred from the normoxic intestine lumen (<5.7% O₂ content) to the more oxygenated tissue (>5.7% O₂). We have seen in the present work that tissue hypoxia damages the oxygen-sensitive germline cells but not somatic cells, which tolerate oxygen and form tumors. It is further conceivable that the GST process also causes an increase in tumor surface antigens (somatic antigens). Similarly, Entamoeba has about 20 different somatic surface antigens, some of which — like cysteine peptidases (lysosomal cysteine peptidases, cysteine cathepsins) and serine-rich proteins (anchored serine proteases) — are potentially deep homologous antigen-sites and archetypes of tumor surface antigens. It will be of interest to learn more about the deep homology of early surface antigens (EAS) in pre-tumoral or early tumoral CSCs and somatic cancer cells; also what changes in EAS architecture occur during germ to soma conversion and the transition from hypoxia to hyperoxia. If deep homologous EAS could be discovered, they could also be targets for vaccines. More about the homologous life cycle of Entamoeba and the amoeba-cancer model occurs separately.

Conflict of interests

The author declares no conflict of interests.

Acknowledgements

I express my gratitude to Dr. Dennis Thomas from the Cold Spring Harbor Laboratory, USA (native English speaker) for reading the manuscript and improving my English. Special thanks to Mr. Gregor Jaruga for the graphics.

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