Cancer: We Should Not Forget The Past

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Abstract

Cancer has been in existence longer than human beings, and man has been facing the illness ever since he made his appearance on Earth. Amazingly, the first human cancer gene was cloned only thirty years ago. This, and other extraordinary scientific goals achieved by molecular cancer research in the last 30 years, seems to suggest that definitive answers and solutions to this severe disease have been finally found. This was not the case, as cancer still remains to be defeated. To do so, cancer must be first understood.

This review highlights how cancer onset and progression has been tackled from ancient times to present day. Old theories and achievements have provided the pillars of cancer understanding, laying the basis of ‘modern era’ cancer research, are discussed. The review highlights the discovery of oncogenes and suppressor tumor genes, underlining the crucial role of these achievements in cancer diagnosis and therapies. Finally, an overview of how the modern technologies have given impetus to expedite these goals is also considered.

Key words: Cancer theories, oncogenes, genomics.

Introduction

The scientific path taken to understand cancer is paved with thousands of theories and scientific findings, from the time of Hippocrates to genomics today. Many of these are fundamental in our modern approaches to the disease.

Chemical carcinogenesis was brought to light with the observations of Paracelsus on cancer onset and environmental exposure to chemicals undertaken over six centuries ago (1-3). The relationship between cancer and chemicals captured the interest of scientists since the 18th century, and this having had a greater impact in the last century. This knowledge, previously unproved, is a reality in today’s research and plays a vital role in the understanding of cancer.

Exactly a century ago, in the book, *Zur Frage der Entstehung Maligner Tumoren*, Boveri presented his theory on cancer that was based on chromosomal abnormality (4-6).

Boveri had no knowledge on the chromosome structure; however, his theory today is considered a pillar of modern molecular approaches to cancer. The discovery of the DNA structure and the achievements of molecular biology have reaffirmed Boveri’s theory and given technological support to developments in this field.

Without doubt, the first human transforming gene, and more broadly, oncogenes and anti-oncogenes, had been previously theorized by Theodor Boveri.

Nowadays, even though scientific research moves ahead rapidly, cancer still needs solid and reliable solutions. Nevertheless, science should look back to historical achievements in advancing new theories and data. Considering the past, and reviewing the milestones of every scientific finding could help in molding future experimental approaches in cancer research.
From Pre-History to Boveri

Cancer has been in existence on Earth even before the appearance of man, as evidenced by paleontological findings of tumors in animals (1, 2).

The first description of human cancer can be found in the Edwin Smith Papyrus dated 3000 BC that illustrated a case of breast cancer. Other documented proof includes the Ebers Papyrus dating from 1500 BC that describes several types of tumors concerning skin, uterus, stomach, and rectum. These old Egyptian documents recorded cancer as a grave incurable disease and associated it to ‘the curse of the gods’ (1). Interestingly, this belief continued to be accepted right up to Hippocrates (460-370 BC), who postulated the earliest scientific theory about cancer. Hippocrates hypothesized that this disease was correlated to an excess of black bile. He believed that cancers, and more broadly any disease, developed whenever the balance in the four body humours (blood, phlegm, yellow bile, and black bile) were lost. Whenever black bile became the dominant humour in some part of the body, this developed cancer.

Hippocrates theory established the foundations of experimental approaches in understanding and confronting cancer, thereafter practiced by all scientists. Claudius Galen (130-200), a Greek physician who practiced medicine in Rome, implemented the Hippocrates theory. He proposed that black bile caused incurable cancer, whereas yellow bile caused curable cancer.

Additionally, the terms and words used to name this disease, and whatever it was associated with, were created by these early ‘scientists’. Hippocrates used the word carcinoma, comparing the disease to a crab (Karkinos) that adheres to its surroundings with its claws (1, 2). The physician Celsus (25 BC-50), later translated this word into cancer, the Latin word for crab. Galen (130-200) described tumors using the Greek term for swelling, oncos.

In the Middle Ages, religious beliefs inhibited the advancement of knowledge, and people began to believe that cancer was an infectious disease.

Eventually, medical science gained a foothold during the Renaissance when scientists began performing autopsies, combining the study of the human body and that of cancer, leading to important achievements. In the 16th century, the anatomist Andreas Vesalius and others were able to demonstrate the non-existence of black bile. Nevertheless, Hippocrates’ theory still continued to have great number of supporters, and it took some time before it was superseded by a new hypotheses. In the same century, Paracelsus studied tumors of mine workers, and suggested there were deposits of sulfur and arsenic salts in blood of these workers, causing their cancers (1-3). At this time it was not known how the elements of the environment caused disease; however, this observation pre-empted the future of chemical carcinogenesis research, being the first association between the work environment and cancer.

A century later, Boerhaave suggested, ‘...that cancer was most likely induced by viruses present in water or in soil. Once acquired, the cancer viruses remained in the body, and they could be transferred by contagious infections or by heredity’. This theory was by no means new, as people during Middle Ages understood that cancer was a contagious disease. Moreover, at that time the term virus indicated some kind of toxic substance, therefore in this respect, Boerhaave’s hypothesis was similar to that of Paracelsus. However, Boerhaave first introduced the new concept that cancer could be hereditary (1, 2).

In the 18th century the Italian pathologist, G.B. Morgagni, founded scientific oncology by performing autopsies after death to identify the patient’s disease. Consequently, he reported that cancer was the result of an ‘organ lesion’.

Other scientists and surgeons suggested cancer was a destructive growth of the organ, caused by their internal structural transformation, or derived from coagulated lymph (1, 2).

During the 18th century, different physicians expanded the hypothesis about the association between cancers and chemical exposure. Their observations had recalled the Paracelsus hypothesis, and this established the new and modern approach of cancer epidemiology. John Hill associated the use of snuff with nasal polyps. Bernardino Ramazzini associated breast cancer with reproductive factors (1, 2). Interestingly, in 1775 Percivall Pott noticed that chimney sweepers frequently developed cancer of the scrotum in later life, and suggested that soot was the cause of this type of cancer. In the last decades of the 19th century, Ludwig Rehn, observed an association between exposure to the aniline dye and bladder cancer in industrial workers (2, 3).

The use of the microscope represented an opportunity for researchers to investigate tumors in great detail. Rudolf Virchow stated that ‘cancer is a disease of cells’, and David Hansemann proposed, ‘The cell of the malignant tumor is a cell with a certain abnormal chromatin content’ (2, 4, 6).

At the beginning of the 20th century, several researchers experimentally confirmed that specific chemicals in the workplace and environment were correlated with the carcinogenic process, demonstrating their multistage and multifactor nature. Katsusaburo Yamagiwa and Koichi Ichikawa, in 1915, were able to induce cancer in rabbits by applying coal tar to the skin of these laboratory animals (3). Addi-
tionally, at this time Theodor Boveri had expanded the Hansemann theory. It is worthwhile to mention what Boveri wrote, ‘tumor growth is based on … a particular, incorrect chromosome combination which is the cause of the abnormal growth characteristics passed on to daughter cells’ (4-6). Figure 1 shows the original book cover published in 1914.

A major milestone for modern era cancer was the discovery of the first tumor viruses (7, 8) in chickens by Ellerman and Bang (1908) and Peyton Rous (1911). These and other important achievements on cancer research, from prehistoric times up to present day are summarized by date in Table 1. Of important note, in the beginning of 1900 external factors such chemical; radiation and viruses were accepted as etiological agents of cancer (3-8). Amazingly, both chemical and viral carcinogenesis produced the same results (9), since, independently, they provided evidence that tumors are correlated to mutated genes.

From Boveri to Oncogenes and Anti-Oncogenes

Ellerman and Bang in 1908, induced erythro-myeloblastic leukemia in healthy chickens by an acellular filtrate collected from diseased chickens (7, 8), which later proved to be an RNA virus. In 1910, Peyton Rous isolated a filterable agent from the sarcoma in the breast muscle of a Plymouth Rock hen. This agent was later named the Rous Sarcoma Virus (RSV) and was regarded as the first RNA tumor virus, as in the early 1900’s leukemia was not considered a cancer.

However, the scientific world at this time was skeptical and did not note the importance of these developments. Fortunately, some researchers had persevered in this field achieving some important goals. For instance in 1936, R. Bittner demonstrated that the mammary carcinoma of C3H mice was induced by viruses and this observation was fundamental in RNA tumor virus research (7, 8). Later, on 1953, L. Gross isolated the first virus inducing leukemia in mice (7, 8).

It is worthwhile to mention that in 1907 the Italian researcher, G. Ciuffo, demonstrated the transmission of human warts by inoculating himself with a cellular filtrate (7, 8). Later, these viral particles were characterized as a DNA virus. The significance of this finding was understood only many years later, when the association between cancer and some types of these viruses was demonstrated, currently known as papillomaviruses. DNA tumor virus research and more broadly of other human tumor viruses, will not be discussed in this review; however, in Table 1 the principal achievements in these fields are listed.

Cell culture techniques implementation and the definition of DNA structure have been crucial in determining infections and transformation mechanisms of these tumor viruses, permitting the isolation and characterizations of mutants (10-12). Table 2 shows the achievements on molecular biology and biochemistry from the end of 1800, up to the advent of the human genome. If Table 1 and Table 2 are compared, it becomes quite evident that the deeper knowledge of cellular molecular processes and the implementation of new experimental approaches were decisive in understanding present day tumor genetics. Figure 2 summarises the major achievements obtained in molecular cancer research and molecular biology.

Figure 1. Zur Frage der Entstehung Maligner Tumoren (1914) book cover, kindly provided by Bi.Ge.A.-Bologna University.
Table 1: TIMELINE OF THEORIES AND ACHIEVEMENTS ON CANCER.

| Period | Tumors in prehistoric animals. |
|--------|--------------------------------|
| 5000 BC| Edwin Smith Papyrus is the earliest written description of cancer. |
| 1500 BC| Ebers Papyrus describes cancers of the skin, uterus, stomach, and rectum. |
| 400 BC | Hippocrates proposes the humoral theory: ‘cancer caused by a surplus of black bile in the body’. |
| 25 BC–50 | Aulus Cornelius describes several varieties of cancers in the book, De Medicina. He points out that breast cancers cause death by spreading to distant organs, and recommends surgical therapy. |
| 23–79 | Pliny describes remedies for cancer in the book, Materia Medica. |
| 81–138 | Aretaeus gives the first description and treatment of cancer of the uterus. |
| 100–200 | Claudius Galen (130–200) implements Hippocrates’ concepts. He proposes that thick black bile causes incurable cancer, and thin yellow bile causes curable cancer. Galen uses the Greek word melancholy (swelling) to describe tumors. |
| 300–400 | Oribasius of Baghdad (325–403) confirms that tumors are formed by black bile. |
| 500–1300 | Aetius (Constantinople 527–565) introduces treatment of breast cancer by amputation of the whole breast. |
| 1100 | Several scientists propose that cancer is caused by ‘germs’, and metastasis was explained as parasites attacking consecutive organs. |
| 1215 | The use of microscopes and the implementation of histological techniques permits the analysis of anomalous cell nuclei in tumors. |
| 1500s | Paracelsus hypothesizes that cancer is caused by accumulation of harmful substances in the bloodstream. |
| 1600s | A. Pare, a French surgeon, hypothesises the cause of cancer as being attributed to an irregular diet that induced an accumulation of feculent material in the blood. |
| 1700s | Physicians propose cancer caused by coagulation and fermentation of blood or lymph. Possible causes of cancer include chronic inflammation, injuries, traumas and familial predispositions. Autopsies highlight cancer as an ‘organ lesion’. |
| 1713 | B. Ramazzini notices the virtual absence of cervical cancer and high incidence of breast cancer among nuns. The scientist associates the different tumor incidences to lifestyle, namely sexual abstinence. This is the start of ‘cancer epidemiology’. |
| 1750 | G. B. Morgagni suggests that cancer is correlated to pathological lesions of a particular organ. |
| 1838 | J. Bittner discovers the mouse mammary RNA tumor virus, transmitted among laboratory strains of mice by breast feeding. |
| 1889 | S. Paget proposes the theory of ‘seed and soil’, to explain metastasis formation. |
| 1877 | J. Cohenheim suggests that tumors arise from embryonic cells. |
| 1860 | R. Virchow proposes cancer as disease of cells. |
| 1907 | G. C. Claudio demonstrates that human warts are caused by an infectious agent. |
| 1914 | T. Boveri hypothesises: ‘tumor growth is based on …a particular, incorrect chromosome’. |
| 1928 | J. Bittner discovers the mouse mammary RNA tumor virus, transmitted among laboratory strains of mice by breast feeding. |
| 1933 | R. Shope discovers cottontail rabbit papillomavirus or Shope papillomavirus, the first mammalian DNA tumor virus. |
| 1936 | J. Bittner discovers the mouse mammary RNA tumor virus, transmitted among laboratory strains of mice by breast feeding. |
| 1937 | A. Graff, C. Friend, and J. Moloney, independently report the identification of mouse leukemia RNA viruses. |
| 1945 | L. Gross isolates the murine polyoma virus. |
| 1957 | B. Eddy and M. Hillman and B. Sweet discover independently the Simian Vacuolating virus 40. |
| 1961 | A. Epstein, B. Achong and Y. Barr identify the first human cancer virus, the Epstein-Barr Virus (EBV). |
| 1964 | H. Temin proposes the theory of provirus hypothesis. |
| 1967 | B. Blumberg isolates and characterizes Hepatitis B virus. |
| 1969 | R. Huebner and G. Todaro propose the vireogene/oncogene hypothesis. |
| 1970 | H. Temin and D. Baltimore, individually, isolate the reverse transcriptase. |
| 1973 | J. Rowley demonstrates chromosome abnormalities in cancer patient. |
| 1974 | A. Levine, D. Lane and L. Old individually discover p53 anti-oncogene. |
| 1975 | discovery of Human T-lymphotropic virus 1 (HTLV I), the first human retrovirus. |
| 1978 | discovery of the first human transforming genes is characterized. |
| 1980 | discovery of Human Papilloma Viruses 16 and 18 are discovered. |
| 1981 | discovery of Rb anti-oncogene from Retinoblastoma. |
| 1982 | discovery of NF2 from Schwannomas and meningiomas. |
| 1985 | discovery of NF1 anti-oncogene from Neurofibromatosis type 1. |
| 1986 | discovery of BRCA1 and BRCA2 anti-oncogenes from breast cancers. |
| 1990 | discovery of NF1 anti-oncogene from Neurofibromatosis type 1. |
| 1991 | discovery of NF2 from Schwannomas and meningiomas. |
| 1992 | Low Resolution Linkage Map of entire human genome published. |
| 1994 | discovery of Kaposi sarcoma-associated herpesvirus (KSHV or HHV8). |
| 1996 | discovery of BRCA1 and BRCA2 anti-oncogenes from breast cancers. |
| 1998 | discovery of Kaposi sarcoma-associated herpesvirus (KSHV or HHV8). |
| 2003 | discovery of Wt1 anti-oncogene from Wilms’ tumor. |
| 2005 | discovery of Nf1 anti-oncogene from Neurofibromatosis type 1. |
| 2008 | discovery of Rb anti-oncogene from Retinoblastoma. |
| 2015 | discovery of BRCA1 and BRCA2 anti-oncogenes from breast cancers. |
| 2018 | discovery of Kaposi sarcoma-associated herpesvirus (KSHV or HHV8). |
Table 2: MOLECULAR BIOLOGY TIMELINE UP TO THE HUMAN GENOME PROJECT.

| Year | Event |
|------|-------|
| 1848 | W. Hofmeister observes chromosomes separation during cell division. |
| 1850 | T. Schwann and M. Schleiden propose independently the cell theory. |
| 1859 | C. Darwin publishes the book, *On the Origin of Species*. |
| 1865 | G. Mendel performs hybridization experiments in pea plants, and proposes the theory of segregation and independent assortment and coined the terms dominant and recessive traits. |
| 1869 | F. Miescher localizes DNA in the cellular nucleus, naming it ‘nucleic acid’. |
| 1875 | C. Darwin introduces the concept of ‘germcells’ as mechanism of inheritance. |
| 1885 | W. Roux formulates the hypothesis of chromosomes as the carriers of inheritance. |
| 1900 | H.M de Vries, proposes the theory of ‘pangenes’ in germinal cells, and introduces the term ‘mutation’. |
| 1902 | A. Garrod discovers the first genetic disease. |
| 1907 | W. Sutton and T. Boveri propose the chromosome theory. |
| 1908 | W. Bateson and R. Crundell Punnett show that the actions of some genes modify actions of other genes. |
| 1911 | A. Sturtevant constructs the first genetic map. |
| 1914 | T. Morgan, and then C. Bridges, demonstrate that genes are carried on chromosomes. |
| 1926 | T. Morgan publishes the book, *Theory of the gene* based on Mendelian genetics and demonstrates that X-ray can induce mutations. |
| 1933 | A. Tiselius introduces the electrophoresis technique for separating proteins in solution. |
| 1937 | F. Bawden discovers tobacco mosaic virus RNA. |
| 1941 | G. Beadle, and E. Tatum propose the one gene concept. |
| 1944 | O. Avery, C. Mcleod and M. McCarty suggest that genes are made of DNA. |
| 1946 | E. Schrödinger publishes the book *What is Life?*, suggesting a complex molecule with the genetic code for living organisms. |
| 1946 | B. McClintock reports the ‘jumping genes’, named transposable elements. |
| 1946 | E. Lederberg discovers the bacteria conjugation. |
| 1949 | M. Delbruck and A. Hershey discover the genetic recombination. |
| 1952 | R. Franklin and M. Wilkins perform X-ray crystallography studies of DNA. |
| 1953 | J. Watson and F. Crick propose the double-stranded, helical, complementary, anti-parallel model for DNA. |
| 1955 | S. Brenner, F. Jacob and M. Meselson discover messenger RNA. |
| 1956 | T. Morgan, and then C. Bridges, demonstrate that genes are carried on chromosomes. |
| 1957 | W. Sutton and T. Boveri propose the chromosome theory. |
| 1958 | M. Nirenberg and H. Gobind Khorana complete the genetic code. |
| 1959 | F. Jacob and J. Monod discover the ‘operon’. |
| 1960 | M. Nirenberg, H. Matthaei and S. Ochoa determine the ‘genetic code’. |
| 1966 | M. Nirenberg and H. Gobind Khorana complete the genetic code. |
| 1967 | M. Weiss and H. Green develop the somatic cell hybridisation. |
| 1968 | W. Fitch and E. Margoliash propose the first evolutionary trees from protein sequences. |
| 1970 | H. Temin and D. Baltimore, independently, isolate reverse transcriptase. |
| 1971 | P. Berg makes the first recombination DNA in ‘vitro’. |
| 1973 | H. Boyer and S. Cohen use a plasmid to clone DNA. |
| 1973 | S. Cohen and H. Boyer report the construction of functional organisms. |
| 1975 | G. Kohler and C. Milstein produce the first ‘monoclonal antibodies’. |
| 1976 | E. Southern publishes the data on Southern Blot technique to identify DNA fragments. |
| 1977 | W. Gilbert and F. Sanger develop the technique to determine the sequence of bases in DNA. |
| 1978 | D. Botstein develops the method ‘restriction fragment length polymorphisms’. |
| 1981 | K. Mullis develops the polymerase chain reaction (PCR). |
| 1984 | A. Jeffreys develops ‘genetic fingerprinting’. |
| 1985 | FASTP/FASTN software introduces algorithms sequence similarity searching. |
| 1986 | Applied Biosystems introduces the first automated DNA fluorescence sequencer. |
| 1987 | D. Burke invents the ‘Yeast artificial chromosomes’, YACs. |
In 1961, RSV was shown to contain an RNA genome (13), and it became the prototype, ‘RNA tumor virus’ (14). Moreover, experiments performed by Temin with actinomycin and amethopterin, inhibitors of nucleic acid, suggested that the virus replicated by transferring the genetic information of the infecting virus, from RNA to DNA, and back again to the RNA of the progeny virus (15).

These results have to be considered as a milestone in tumor virology, although the RNA genome of the virus represented a dilemma hard to resolve.

In 1964, Temin proposed the theory of provirus, ‘...virus enters a cell and directs formation of a DNA containing the genetic information of the virus. This new DNA, the provirus, then acts as a template for the formation of new nucleic RNA, for the virion....’ (16).

The provirus theory was based on the revolutionary hypothesis that RSV replicates through a synthesized DNA intermediate using as a template the RNA genome (17). Up to this time, no evidence existed on the possibility of transferring genetic information from RNA to DNA and back to RNA. Hence, this innovative model of viral replication did not receive much attention from the scientific community. However, researchers had the urgency to resolve the inexplicable results on RNA tumor virus replication obtained with inhibitors of nucleic acid synthesis. In 1969, Huebner and Todaro proposed the virogene/oncogene hypothesis. This theory postulated that DNA of most, or all of, the eukaryotic cells carried vertically transmitted RNA tumor virus information, this being known as virogene (18). These viral sequences contain the oncogene, which may transform normal cells into tumor cells. These RNA tumor viruses are present in a repressed form in the cells and they may be ‘switched on’ by various environmental factors, and also are dependent on the host genotype. The ‘switching on’ of these sequences can produce an infectious virus and/or induce the cellular transformation. The virogene hypothesis differs from the provirus hypothesis, since the latter implies that an exogenous virus infects the cell. Instead, the virogene/oncogene is part of the genetic background of all vertebrate cells. In 1970, Temin and Baltimore individually isolated the reverse transcriptase, which was able to mediate the RNA-dependent DNA synthesis (19, 20). This enzyme represented the key element of the provirus hypothesis, and after this event the RNA tumor viruses were renamed retroviruses.

Reverse transcriptase had definitively changed the approach in studying RNA tumor viruses and permitted new experimental methods in investigating their neoplastic mechanisms. In 1976, Dominique Stehelin, Harold Varmus, Michael Bishop, and Peter
Vogt characterized the sequences of RSV responsible for its neoplastic potential, the src gene (21). Moreover, they demonstrated the cellular origin of this transforming gene, since its normal counterpart is present in the normal eukaryotic cells. Following this, many more transforming genes of other tumorigenic retroviruses were characterized, and their cellular origin confirmed. These achievements recalled the theory of viogene/oncogene by Huebner and Toda-ro, and these genes were actually named oncogenes, and their cellular counterpart proto-oncogene (22).

Around this time, Bruce Ames demonstrated that carcinogens could mutate bacterial genes (23). The experiments proved that carcinogens could damage the genes of target cells, confirming the theory that cancer cells carried mutant cellular genes, which may confer a growth advantage.

The achievements on the transformation mechanisms of retroviruses and the findings on the mutagen activity of carcinogens demonstrated that tumors had gene alteration as a common denominator, independent of their etiology (6, 22, 23).

Researchers had felt the urge to go further, and formulated experimental protocols in defining possible relationships between viral oncogenes and genes mutated by physical and chemical carcinogens.

Early in the 1970s, Graham and van der Eb developed a procedure for gene transfer in mammalian cells (24). The researchers used this method to introduce DNA of virally transformed cells into NIH 3T3 recipient cells, obtaining foci of transformation in their recipient cells several weeks later (25). Hence, this approach appeared to be a suitable method to analyse the genetic background of cancer cells, and to eventually reveal the genes involved in the neoplastic process.

DNA transfection was then used to analyze the presence of transforming active sequences in DNA extracted from several chemically transformed 3-methylcholanthrene (3-MC) mouse-cell lines. This experiment yielded a large number of foci, and non-transforming or activated endogenous viruses being detected in these mouse cells. It was probable that the transforming activity was derived from a cellular gene mutated by the 3-MC into a transforming active allele (25).

Later, researchers used the gene transfer protocol to investigate the presence of active transforming genes in human tumor cell lines and primary tumors. Remarkably, three different research groups demonstrated that DNA extracted from human tumors cells lines could transform NIH/3T3 cells. Furthermore, the characterization of the transforming sequences detected in the cell line T24/EJ, established from a human bladder carcinoma, evidenced their homology to ras, the transforming gene of Harvey, and the Balb sarcoma viruses (Ha-ras) (26-28). Molecular comparisons between the human ras transforming gene and its normal allele revealed that the malignant potential was activated by a single point mutation. Also, the ras gene can acquire a transforming activity from mechanisms other than capture by retroviral sequences (9). Investigators continued with the transfection approach analyzing DNA extracted from human solid and hematopoietic tumors (29, 30). Surprisingly, most of the activated transforming sequences detected in these tumors were homologous to viral ras genes (Ha-ras and Ki-ras), although these later experiments also revealed activated transforming sequences that did not have viral counterparts (30). Moreover, ras transforming sequences were detected also in animal tumors induced by chemical carcinogens (9, 31). These results rolled back again to the Boveri’s concept of neoplastic process in consequence of genetic alterations.

In the mid-1980s, DNA-transfection experiments confirmed the old theories on cancer onset and development, and fitted in well with tumor virology and chemical carcinogenesis. Despite, these extraordinary achievements, investigators soon underlined the limits of viral transforming genes had been used to investigate at molecular level tumors and their cells. Many human sequence homologues of viral oncogenes were cloned and localized in human chromosomes. These molecular approaches highlighted the fact that human proto-oncogenes can acquire transforming activity and drive cells to the neoplastic phenotype if rearranged, translocated or amplified (22).

One of the most relevant examples is the translocated Philadelphia chromosome associated with chronic myelogenous leukemia, a hematopoietic tumor. This translocation involves the Abl gene on chromosome 9 and the BCR (breakpoint cluster region) gene on chromosome 22 (32). Also, Bukitt’s lymphoma cells were shown to present three types of translocations, all involving a fragment from the end of the long arm of chromosome 8, which can be translocated to the long arm of chromosomes 14, or eventually, to chromosomes 2 or 22. The translocated fragment of chromosome 8 contains the myc gene. This myc sequence can be translocated to chromosome 14, near to the immunoglobulin heavy chain locus, or to chromosomes 2 or 22, near to the two immunoglobulin light chain genes (33, 34). Myc gene amplification was found in the human promyelocytic leukaemia line HL60, and other cancer lines (35).

Moreover, it was well known from cytogenetic analysis that tumor cells may show loss of genetic information. Researchers developed sophisticated
protocols based on cell hybrids to investigate at molecular-level these genetic abnormalities. By cell hybridization of normal fibroblasts with the human cancer cells, HeLa non-tumorigenic clones were obtained, some of which reverted to neoplastic phenotype (36). Molecular cytogenetic analysis of these clones had evidenced that the transformed phenotype correlated with the loss of the fibroblast chromosomes 11 and 14. Together, these findings proved that neoplastic phenotype and loss of genetic sequences are correlated. Furthermore, the molecular characterization of retinoblastoma and Wilms’ tumor evidenced deletions on both pairs of chromosomes 13 and 11, respectively (37, 38). Additionally, transfection of a normal allele of chromosomes 11 into tumorigenic Wilms’ tumor cells can revert to their phenotype. The chromosomal locus/genes correlated with these two neoplasia had been cloned and characterized, following elaborate cloning strategies.

The genes, associated with retinoblastoma and Wilms’ tumor development were named Rb1 and Wt1, respectively. Retinoblastoma and Wilms’ tumor are both heritable and non-heritable tumors. Their characterization demonstrated that germ-line mutations are present in the heritable form (22, 37, 38). However, this inherited mutation is not sufficient for tumor formation; a second mutational event is required. Similarly, for the development of non-heritable tumors, two somatic events must occur and inactivate both genes. These results proved the two-hit hypothesis of Knudson according to ‘a complete inactivation of both alleles of a tumor-suppressor gene is required for cancer onset and development. …..’. Hence, these sentinel genes acting against tumors have been named anti-oncogenes or tumor suppressor genes (39, 40).

Characterization of the protein products of proto-oncogenes and tumor suppressor genes revealed their role in regulating intra- and extracellular signaling related to cell growth and division. Usually proto-oncogene products are correlated with cellular functions that stimulate cell growth and division. Tumor suppressor genes are normal genes, mainly devoted to the cellular processes correlated with cell growth inhibition and programmed cell death (apoptosis) (6, 22, 33).

Mutation, amplification, deletion or translocation of these genes disrupt their function, leading to an uncontrollable growth and cancer (6, 22).

Moreover, epigenetic changes also contribute to carcinogenesis. These changes alter the DNA and associated proteins without changing the DNA sequence, modulating the chromatin structure and DNA accessibility. Through these mechanisms, epigenetic changes may alter the transcriptional status of cancer genes, or regions of chromosomes that control their expression, inducing the neoplastic process (41). Interestingly, anti-oncogenes are involved in tumor development induced by some DNA tumor viruses. These viruses code proteins that are capable of inhibiting tumor suppressor proteins. An example, is the binding between p53 protein and viral transforming proteins such E6 of oncogenic HPV (40, 42).

Boveri had predicted all these findings, when he wrote, ‘…Another possibility is that there is a specific inhibitory mechanism in every normal cell that only permits cell division to take place when this mechanism is overcome by some special stimulus…A tumor cell that proliferated without restraint would be generated if these ‘inhibitory chromosomes’ were eliminated’ (4, 5).

Past, Present and Future

In 1914, Boveri wrote, ‘It is conceivable that for any one cell type there is one particular abnormal combination of chromosomes that endows the cell with the properties of malignancy…’, predicting cancer to be a complex multifaceted process. Today, it is understood that more than one gene/chromosomal alteration contributes to tumor development, and these must be known and characterized to fully comprehend the molecular process of tumor development.

In 1985, Dulbecco stated in support of these factors, ‘A major gap in our understanding of cancer is how the activity of an oncogene is related to the events of progression. But, the first task is to ascertain whether the DNA of an advanced cancer is as heterogeneous as the phenotype of its cells. If we wish to learn more about cancer, we must now concentrate on the cellular genome……..’, and in consequence, he advocated the Human Genome Project (HGP) (43). As Dulbecco had foreseen, the sequencing of the human genome represented the corner stone for the construction of physical and genetic maps and the identification of genes involved in human cancer.

Integrating the knowledge on human genomes with the new powerful technologies for DNA sequencing and inspection of protein expressions (mass-spectrometry), has permitted the collection of large amounts of data on genetic alterations and aberrant gene expression correlated to onset and development of different cancers (44, 45). The Cancer Genome Atlas (TCGA, http://cancergenome.nih.gov/) and the International Cancer Genome Consortium (ICGC, https://icgc.org/) provide clinical data and the genomic profiles of thousands of normal and tumor samples for a variety of cancer types. This has enabled the identification of many novel cancer genes. The catalogue - that includes oncogenes, anti-
oncogenes, epigenetic regulators, cell cycle-associated genes, DNA damage response/repair genes, metabolic regulators, and microRNAs - has not yet been completed (46-49). Cellular and genomic data from a variety of tumors have confirmed and highlighted their extraordinary heterogeneity. This heterogeneity has been evidenced among tumors from patients with the same cancer type, inter-patients heterogeneity, and within cancer cells of an individual patient's tumor, intra-tumor heterogeneity. Heterogeneity represents, indeed, a thorny challenge to overcome in order to provide clinics with a suitable cancer genomic data. This objective does not appear simple to attain. The inter-patient heterogeneity is most likely tied to the different genetic backgrounds and lifestyles, and they should be considered and evaluated. Therefore, a molecular investigation at single-patient level may be required for a correct diagnosis and optimal therapy (48, 49). Moreover, intra-tumor heterogeneity should also be taken into account, and the mechanisms producing it. To explain the intra-tumor heterogeneity, one of the proposed models is the 'clonal evolution model'. It assumes that different mutant clones acquire a survival advantage, because of the natural competition with other clones. Stochastic genetic or epigenetic changes are claimed as the most probable mechanisms conferring growth advantage to cancer cells. Likely, macro and micro-environments play an important role in the development of the subpopulations of cancer cells, enabling them to initiate and/or propagate the disease, or even to permit its remission (50, 51).

Recently, cancer stem cells (CSCs) have gained the interest of scientists as a good candidate and 'model' to investigate tumor heterogeneity. Much experimental evidence has highlighted the presence of CSCs in a broad spectrum of malignancies. These cells can self-renew, proliferate and differentiate. It has been suggested that some tumors may arise from small populations of CSCs that give rise to phenotypically diverse cancer cells, playing important roles in tumor heterogeneity, favoring growth, dormancy or recurrence, and metastasis (52). The CSCs imply that tumors are organized as an adult tissue, and their genetic heterogeneity of cancer propagating cells may derived from 'clonal evolution' within the stem cell pool by the Darwin’s premise of 'natural selection'. These models of cancer initiation and spreading refers back to the ‘seed and soil’ and ‘embryonal’ theories proposed by Paget and Cohenheim, respectively, more than hundred years ago (Table 1). Moreover, they emphasize that a tumor is composed of complex mixtures of cells with various genetic alterations and functions. They then suggest a single-cell analysis as feasible approach to overcome the dubious evaluation caused by multiple mixtures of mutations within tumor cells. Nowadays, single-cell genomic sequencing and single-cell transcriptomic sequencing have been explored as potential techniques to investigate cancer heterogeneity. Single-cell genomic sequencing mainly focuses on the general landscape of mutations, such as single nucleotide variations, insertions, and deletions in a single cell (53). Single-cell transcriptomic sequencing can be useful for analyzing genetic network regulation, including differentiation, reprogramming and transdifferentiation. Single-cell transcriptomic sequencing can highlight transcriptomic alterations (mRNAs, microRNAs, retained introns, alternative splicings, long-noncoding RNAs and fusion genes) (50-53). Data coming from such analysis could be relevant in understanding the role and the fate of single cells in a tumor, mainly CSCs. These techniques, along with appropriate ‘in vitro’ and ‘in vivo’ CSCs models to mimic tumor growth, could give an insight into genetic alteration/modification. It could also provide an understanding of the gene regulation networks responsible for physiological functions, behaviors, and phenotypes in response to signals and microenvironmental changes. The incoming data could be useful for revising and making available the huge amount of genomics information for the world research community. Moreover, this information, when cross-referenced and integrated, could be crucial to define better personalized therapeutic strategies (48-53), as well as to design a broad spectrum of preventive and therapeutic strategies that are broadly relevant. The analysis of expressed biomarkers in the cancer is one of the most used approaches for the disease’s treatment, and recently, particular attention has been given to CSCs, as a main factor in fighting cancer. Different therapeutic strategies have been developed and are being designed to attack the CSCs and to destroy their surrounding environment; some of these strategies are under preclinical and clinical evaluation. Some strategies rely on using monoclonal antibodies against surface markers that are different from normal stem cells. Monoclonal antibodies against these markers and conjugated with cytotoxic agents permit the targeting of the CSCs cells and eventually kill them.

CSCs cells play also an important role in chemo-resistance acquisition. To overcome this problem, the mechanism responsible for the chemo-resistance must be identified and characterized. One of the most recognized mechanisms is the efflux of cellular cytotoxic drugs by the ATP-binding cassette (ABC) transporter proteins. The trans-membrane proteins using the energy from the ATP hydrolysis expel the drugs from the cells. Studies are focused on molecular drugs capable of inhibiting
the ABC transporter protein. Moreover, CSCs could be attacked modifying the specific pathways involved in apoptosis and resulting in suppression of cellular characteristics. Particular attention has to be given to the tumor micro-environment, since it plays a relevant role in CSCs growth and selection, as well as in protecting them from drugs. For instance, experimental evidence shows a protective role of bone stromal cells in B cell malignancies. Finally, several therapeutic approaches are focusing on the angiogenesis processes as a mechanism to control CSCs growth and differentiation.

A different strategy is to target genes that are affected in a high percentage of tumor types, such as Myc or Ras, and to which cancer cells appear to be ‘addicted’ (54, 55). Approximately 30% of human tumors harbor a mutant Ras gene, with an altered GTPase activity. Targeting the catalytic domain of the Ras proteins is technically challenging. Therefore, many efforts have been directed at alternative strategies such as the synthetic lethal approach, which aims at targeting proteins that are crucial for the survival of tumor cells with an activated Ras protein. Theoretically, it may be possible to kill tumorigenic cells expressing a mutated Ras gene without affecting normal cells (55).

These are just few examples of the recent efforts to relocate the modern achievements in cancer research into the diagnosis and therapy fields. However, much more is still to be done. More genomics data, more animal-models that are suitable, more implemented personalized medicine trials, and new expertise such as onco-bio-informatics, will be crucial to implement onco-knowledge into clinical evaluation. Moreover, scientists should always consider the ethical, legal, and social implications that clinical-genomics raise. For instance, the high costs of such therapeutic and diagnostic approaches that can put a financial strain on people selecting treatments that they can or cannot afford. This, in the end, could determine who may or may not undergo the most appropriate cure.

The purpose of this review is to underline past developments that may help to achieve important new goals in cancer research. Interpreting and construing the past giants theories relating to the modern achievements have been a quite exciting, but easy task. However, predicting the future of cancer research, which is based on current and past knowledge, is difficult and should take in account two considerations. The first is the speed of developments: nowadays research moves so fast. The second is the complexity: the modern technologies generate thousands of results, and each one is a small element looking for its place in an extraordinary puzzle. Therefore, these peculiarities of current cancer research suggest that future investigations should be based on worldwide cooperation, the best ‘technology’ for reaching the daring goals to understand cancer and ‘eventually’ cure it.

Conclusions

Cancer research evolved down a long path, since Hippocrates proposed his theory of black bile. Bo- veri’s vision of cancer as a genetic disease a century ago, cloning of the first human oncogene almost thirty years ago, and decoding of the human genome almost ten years ago, represent major steps in the understanding of cancer. Nowadays, DNA sequencing and gene expression profiles may have a relevant prognostic value in a wide variety of malignancies. For instance, DNA copy number variations and gene expression signatures are employed to define tumor subtypes and as prognostic indicators in chronic myeloid leukaemia, breast cancers and other tumors (56, 57). Bioinformatics technologies and cancer databases are used to combine molecular, clinical, and population data for diagnostic purposes and for formulating therapeutic strategies. By analyzing individual genetic profiles, it may be possible to foresee risks of cancer development and devise personalized therapies (58, 59). We are now in the era of cancer ‘omics’, which investigate genomes, epigenomes, methylomes, transcriptomes, miRnomes, metabolomes, proteomes and so on, easily accumulating ponderous amounts of information, often causing intractable problems (46). There are still huge gaps among these large datasets and in the understanding of cancer biology. How the cancer genomes and gene expression programs interact to dictate cancer cell behaviour, the role of cancer stem cells, the mutual influence of the cancer cells with the cells of the microenvironment, are all aspects that have to be confronted.

Still cancer is an unresolved problem, and researchers must work harder to get to the root of all these open ended questions.

Confucio said, ‘Study the past if you would define the future’. Past scientific achievements could represent the milestones for building future bridges along the path of cancer research understanding, ‘we should not forget the past’.

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Competing Interests

The authors have declared that no competing interest exists.

References

Hajdu SI. A note from history: landmarks in history of cancer, Part I. Cancer. 2011; 117:1097-102. doi: 10.1002/cncr.25533.

Mitsui I, Bryndza E, Soochani A, Scala S. Evolving models of tumor origin and progression. Tumor Biol. 2012; 33:914-7. doi: 10.1007/s13277-012-0389-0.

Henschel F. Yamagawia’s tumour and its historical significance. From Percival Pott to Katsushibama Yamagawa. Gann. 1968; 59:447-51.

Boveri T. Zur Frage der entstehung maligner tumoren. Jena, Germany: Gustav Fischer; 1914.

Boveri T. Concerning the origin of malignant tumours by Theodor Boveri. Translated and annotated by Henry Harris. J Cell Sci. 2008; 121 (Suppl 1): 1-84. doi: 10.1242/jcs.025742.

Baltimore A. Cancer genetics: from Boveri and Mende to microarrays. Nat Rev Cancer. 2001; 1:177-82. doi: 10.1038/3504086.

Javier RT, Butel JS. The history of tumor virology. Cancer Res. 2008; 68:7693-706. doi: 10.1158/0008-5472.CAN-08-3301.

Kalland KH, Xi Song Ke XS, Oyan AM. Tumour virology - history, status and future challenges. APMS. 2009; 117:382-99. doi: 10.1111/j.1600-0463.2009.02452.x.

Barbacid M. ras genes. Annu Rev Biochem. 1987; 56:821-48.

Watson JD, Crick FH. Molecular structure of nucleic acids. Nature. 1953; 171:730-3.

Temin HM, Rubin H. Characteristics of an assay for Rous sarcoma virus and Rous sarcoma cells in tissue culture. Virology. 1958; 6:69-68.

Water JD, Crick FH. Molecular structure of nucleic acids; a structure for deoxyribose nucleic acid. Nature. 1953; 171:730-3.

Hollday R. The early years of molecular biology: personal recollections. Notes Rec R Soc Lond. 2003; 57:195-208. doi: 10.1098/rpr.2003.0206.

Crawford LV, Crawford EM. The properties of Rous sarcoma virus purified by density gradient centrifugation. Virology. 1961; 13:227-32.

Drayton HA. The inactivation by organic solvents and detergents of partially purified Rous I virus preparations. Br J Cancer. 1961; 15:348-53.

Tjem HM. The effects of actinomycin D on growth of Rous sarcoma virus in vitro. Virology. 1963; 1:79-83.

Tjem HM. The participation of DNA in Rous sarcoma virus production. Virology. 1964; 23:486-94.

Tjem HM. Malignant transformation in cell cultures. Health Lab Sci 1964; 179-83.

Huebner RJ, Todaro GJ. Oncogenes of RNA tumor viruses as determinants of cancer. Proc Natl Acad Sci USA. 1969; 64:1087-94.

Baltimore D. RNA-dependent DNA polymerase in virions of RNS tumor viruses. Nature. 1970; 226:1209-11.

Tjem HM, Mizutani S. RNA-dependent DNA polymerase in virions of Rous sarcoma virus. Nature. 1970; 226:1211-3.

Steinher D, Varmus HE, Bishop JM, Vogt PK. DNA related to the transforming gene(s) of avian sarcoma viruses is present in normal avian DNA. Nature. 1976; 260:137-3.

Weinberg RA. The molecular basis of oncogenes and tumor suppressor genes. Ann N Y Acad Sci. 1995; 758:331-8. doi: 10.1111/j.1749-6632.1995.tb42838.x.

Amin BN. Mutagenesis and carcinogenesis: endogenous and exogenous factors. Environ Mol Mutagen. 1989; 14 (Suppl 16):566-577.

Graham FL, van der Eb AJ. A new technique for the assay of infectivity of human adenosine 5 DNA. Virology. 1973; 52:456-67.

Weinberg RA. Use of mutation to analyze genetic information and malignant transformation. Biochim Biophys Acta. 1981; 651:25-25.

Parada LF, Tabin CJ, Shih C, Weinberg RA. Human EJ bladder carcinoma oncogene is homologue of Harvey sarcoma virus ras gene. Nature. 1982; 297:474-8.

Santos E, Tronick SR, Aaronsen SA, Pulciani S, Barbacid M. T24 human bladder carcinoma oncogene is an activated form of the normal human homologue of BALB- and Harvey-MVSV-transforming genes. Nature. 1982; 298:343-7.

Fasano O, Taparowsky E, Fiddes J, Wigler M, Goldfarb M. Sequence and structure of the coding region of the human H-ras-1 gene from T24 bladder carcinoma cells. J Mol Appl Genet. 1985; 2:173-80.

Eva A, Tronick SR, Gol RA, Pierce JH, Aaronsen SA. Transforming genes of human hematopoietic tumors: frequent detection of ras-related oncogenes whose activation appears to be independent of tumor phenotype. Proc Natl Acad Sci USA. 1983; 80:501-20.

Pulciani S, Santos E, Lauver AV, Long LG, Aaronsen SA, Barbacid M. Oncogenes in solid human tumours. Nature. 1982; 300:539-42.

Sukumar S, Pulciani S, Doniger J, DiPaolo JA, Evans CH, Zbar B, Barbacid M. A transforming ras gene in tumorogenic guinea pig cell lines initiated by diverse chemical carcinogens. Science. 1984; 223:1197-9.

de Klein A, van Kessel AG, Goldfarb M, Bartram CR, Hagemeyer A, Boetsma D, Spurr NK, Heisterkamp N, Groffen J, Stephenson JR. A cellular oncogene is translocated to the Philadelphia chromosome in chronic myelocytic leukaemia. Nature. 1982; 300:765.

Nasi S, Ciarpaccia R, Tucker R, Rosati J,Soucek L. Making decisions through Myc. FEBS Lett. 2001; 490:153-62. doi: 10.1016/S0014-5793(01)2218-4.

Molyneux EM, Rochford R, Griffin B, Newton R, Jackson G, Menon G, Harrison CJ, Israels T, Bailey S. Burkitt’s lymphoma. Lancet. 2012; 379:1244-4.

Dregger P, Schmid FM, Attarian S, Griesinger C, Granducci G, Hengstrockner N, Hengst K, Eberhard P, Langford J, et al. The evolving role of individualized therapy. J Cancer. 2013; 4:736-43. doi: 10.7150/jca.7735.