Curative therapies in cancer: A perspective

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Abstract
Oncologic therapies for adult tumours such as therapy using overall cell division, DNA repair, angiogenesis, growth factor pathways or metabolic pathways can seldom provide long term curation. In future oncology only a few new therapeutic modalities have a potential of offering long term curation relying on mechanisms such as manipulating the immune system or providing organ or cell specific cell killing with knowledge about the normal cell of origin of the tumour. Correcting DNA mutations and interfering with transcription in tumours may be important on coming treatment modalities.

Introduction
It is not always easy to define the cure of cancer for a patient. Generally, it is possible to mention a cure for a group of similar patients diagnosed with the same disease, tumour stage, and receiving the same therapy when the survival curve for the patients become parallel with an age and sex matched group from the general population (Figure 1). In early years, the curation of tumours was overestimated and was later found to more often mirror the natural course of a tumour regardless of the therapy. This was evident when new regression models for survival such as advanced Cox proportional hazard analyses were employed.

There has been a large increase in incidence of common cancers such as breast cancer, prostate cancer, melanoma and colon cancer not accompanied by a similar increase in tumour mortality (Figure 2 for incidence and mortality for some cancers in Sweden). The increase occurs at least partly before opportunistic or organized screening have started or new life saving therapies have been introduced such as immune therapy for patients with melanoma.

The data instead suggest a massive over diagnosis of cancer rather than gains in therapeutic results. When studying effects from modern therapeutic interventions this need to be carefully assessed.

Many tumour therapies have the main effect of prolonging progression-free survival without affecting overall survival rate. While others have as the main effect the ability to improve life quality without affecting survival. Among recent therapies, a growing disappointment has emerged for many of the targeted cancer therapies that mainly only improve progression-free survival [1].

This should be compared with therapy (in parenthesis) directed to the hallmarks of cancer as described by Hanahan & Weinberg [2]. Each hallmark is presented by the example of a possible therapeutic approach (in parenthesis). The hallmarks of cancer include deregulating cellular energetics (aerobic glycolysis inhibitors), resisting cell death (proapoptotic BH3 mimetics), genome instability and mutation (PARP inhibitors), inducing angiogenesis (inhibitors of VEGF signaling), activating invasion and metastasis (inhibitors of HGF/c-Met), tumor-promoting inflammation (selective anti-inflammatory drugs), enabling replicative immortality (telomerase inhibitors), avoiding immune destruction (immune activating anti-CTLA4 mAb), evading growth suppressors (cycin-dependent kinase inhibitors), and sustaining proliferative signalling (EGFR inhibitors). These hallmarks have with time been refined and developed further [3-5].

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Only a few therapies directed against these hallmarks can as single agents improve survival and be curative like immune checkpoint inhibitors. Investigations are ongoing to find out the role of a therapeutic approach of combining therapy against more than one hallmark [6].

**Hypotheses behind cancer therapies**

This section presents a number of hypotheses regarding mechanisms around different cancer therapies and their relationship with the cure of cancer.

**Dividing cells and deficient DNA repair**

Both radiation and cytostatic therapy have their main effects on dividing cells. However, tumour cells have deficient DNA and cellular repair when compared with normal cells. Therefore, fractionating the therapy is a successful way of increasing the therapeutic ratio between neoplastic and normal tissue. Combining agents to counteract tumour resistance and reduce toxicity are of importance compared with single-agent therapy [7,8]. The curative potential of radiation and cytostatic therapy is, however, limited. Recently, a concept of synthetic lethality to target deficient DNA repair through, for instance, PARP inhibition has been added to the therapeutic arsenal [9].

**Growth factor inhibition/hormone inhibition**

Inhibiting growth factors and hormones cause cell arrest or apoptosis. Some successful examples exist such as anti-oestrogen therapy in breast cancer with oestrogen receptor blockade, aromatase inhibition or LHRH inhibition [10]. Also, progression free survival in prostate cancer is improved by inhibition of testosterone [11]. Included among this therapy category is therapy against GIST tumours with imatinib inhibition of kit, PDGRFA and ABL [12]. Her-2 neu inhibition is discussed later in the manuscript.

**Starvation or caloric restriction**

Caloric restriction in animal studies leads to both a lower cancer incidence and better tumour therapy responses [13]. It is therefore conceivable that the same occurs in the human situation as well. Obese cancer patients, in general, have a worse tumour prognosis. The role of caloric restriction and/or physical activity are studied in large cohorts. Therapy with antimetabolites such as methotrexate and 5-fluorouracil are examples of such studies. Antibiotics such as doxorubicin and mitomycin also belong to antimetabolites.

**Cell differentiation**

Some tumours originate from not fully differentiated tissues somewhat resembling embryonic tissue. Agents that promote tissue differentiation have successfully been used in cancer therapy. The cisplatinol therapy of testicular germ cell tumours belongs to this category [14]. The methotrexate therapy of choriocarcinoma and ATRA (all-trans-retinoic acid therapy) in acute promyelocytic leukaemia are other examples.

**Angiogenesis**

Patients who form more blood vessels supplying the tumour tissue with oxygen and nutrients would likely have worse prognosis [15]. Anti VEGF therapy is an example of this type of therapy.

The above listed therapy options have very limited success from a curative standpoint.

Few therapies are able to improve progression-free survival, overall survival and life quality concurrently. An important exception is the successful therapy with cytostatic in childhood cancer and therapy for Hodgkin's disease, non-Hodgkin lymphoma, some leukaemias, testicular cancer and choriocarcinoma [7,8,16]. Antihormonal therapy of breast cancer and imatinib therapy of GIST tumours also offers long term survival. However, new emerging research results can better predict the curative potential of a therapy also in adult patients. Below are mentioned approaches that have strong potentials of being curative.

**Organ or cell population specific cell killing**

Designing cancer therapy based on specific organ toxicity for the normal tissue of the same organ as the tumour could be very fruitful [17]. A number of observations support the hypothesis that there is, at least, partly a common susceptibility for cancer agents between the cell of origin and the tumour itself. A prerequisite for successful therapy is that the individual can compensate for the organ or cell population loss ordamage. Firstly, surgery either of the primary tumour or as prophylactic operations can be curative or preventive [18-20]. Secondly, antibody therapy directed against Her-2 neu (breast cancer), CD20, and CD30 (non-Hodgkin lymphoma) is also curative [21-23]. The patient must be able to tolerate the organ or cell population loss also from the normal tissue which is the case for the above examples. Also, allogenic stem cell therapy of leukaemia is an example of this approach where stem cells from another individual is given back after leukemic and normal bone marrow of the recipient have been eradicated.

Most kidney cancers originate from tubuli cells. In metastatic kidney cancer, the mushroom poison orellanine, that in accidental poisoning only injures the tubuli cells, has been shown to have anti-tumour effects and a potential of curing metastatic disease [24].

**Immunological self-non-tolerance**

Immune escape occurs because the tumour tissue is very similar to the individual and therefore, a strong enough immune response is not evoked. Resetting the immune system by provoking autoimmunity by drug therapy is also favourable for killing the tumour cells and metastic cells. This is clearly seen by therapy against CTLA-4, PD-1 and PD-L1. Long-term survivors are seen for a number of patients with different tumour types such as melanoma, lung cancer, mismatch repair colon cancer, Hodgkin's disease, ear-nose-throat cancer and kidney cancer [25]. Newly by FDA approved genetically engineered Chimeric antigen receptor (CAR) T cells may also be an important new therapy especially for lymphoma and leukaemia [26].

To understand the biology of a tumour, it is important to determine the cell of origin of the tumour.

**Cell of origin**

The biology of a tumour is at least partly determined by the characteristics of the cell of origin. This was first hypothesized through epidemiological data by us and later confirmed by experimental data by Weinberg's group [27-29]. The cell of origin could be of importance for both the invasiveness and metastatic potential of the tumour [30]. This could be manifested through methylation patterns, driver and suppressor genes present in the tumour.

**Summary**

Organ or cell population specific cell killing and modern immune therapy can offer possibilities for long-term cancer survival/cure. Predicting the biology of a tumour especially as characterized by its normal cell of origin will further aid in combatting its invasiveness and metastatic potential. Some tumours, such as brain tumours and those
with neuro-ectodermal differentiation where little or no therapeutic ratio exists between the tumour and normal tissue will even in the future be hard to treat. Accordingly, tumours from the brain and the nervous system will continue to be hard to treat with the present organ or cell population specific cell killing or immune therapy. The latter is also hampered by not having a similar lymphatic system as the rest of the body.

Improvement and development of gene therapy to reinstitute suppressor genes and/or to inhibit oncogenes (tumour drivers) have in the future a potential role of further improving cancer therapy. CRISPR-Cas9 technology looks especially promising as does therapy interfering with the transcription process of RNA [4,5] (Figure 3).

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