Introduction

Cellular immunogene therapy inducing immune anti-tumor response constitutes one of the latest strategies for the treatment of many forms of cancers. From 2015, the cancer immunotherapy became an obligatory complement therapy in USA: The Cancer Moonshot Task program supervised by USA government and the cancer immunotherapy program established in Parker Institute in Los Angeles—the common program of the best university hospitals in USA [1]. The beginning of our Anti-gene (antisense and triple helix) IGF-I cellular immunogene Phase I trial in USA and Europe has presented promising results: an increase in immune response goes together with life span, and confirms the role of the immune phenomenon in the suppressing of animal tumors treated experimentally by the same cellular immunogene therapy. We need to underline, that our clinical trial was realized following the establishment of immunogene therapy as a clinical domain. This approach is based on suppression of growth factor IGF-I in cancers cells [2-5].

Figure 1: Mechanism of IGF-I antisense immunogene therapy. Cancer cells transfected with a vector encoding IGF-I cDNA in antisense orientation express MHC-I and B7 molecules. These immunogenic cells become also apoptotic. The APC - antigen presenting cells which are involved in tumor-immunogenicity mechanisms activate lymphocytes T CD8 and CD28. The last ones form the bridges with MHC-I and B7 present in the solid tumor inducing anti-tumor processes after every three vaccinations [7,12].

Alteration in expression of growth factors is associated with growth and development of human tumors. One such growth factor is IGF-I, a 70-amino acid polypeptide. Using IGF-I as the diagnosis marker, 17 different tumors are known to express IGF-I gene. IGF-I seems to play a major role in neuroectodermal tumors such as glioblastoma (the most frequent brain tumor in man and usually fatal), as in other tumors - mesodermal i.e. prostate adenocarcinoma, and endodermal i.e. colon adenocarcinoma [6,7]. In the preclinical experimental studies, C6 and CNS-1 rat glioma, or LFC rat hepatoma cells, and PCC3 murine teratocarcinoma cells transfected with a pAnti-IGF-I vector or with a pIGF-I TH vector (antisense or triple helix technology), became immunogenic expressing MHC-I and B7 antigens. Moreover, the transfected tumor cells became apoptotic. These cells injected into syngeneic animals induced a T-cell mediated immune reaction both against themselves and against their non transfected tumorigenic progenitor cells. IGF-I antisense or triple helix transfecants were shown to elicit a curative anti-tumor immune response with tumor regression at distal sites [8] (Figure 1). Our strategy of treatment of malignant tumors is based on:

a) Diagnosis using IGF-I gene expression as differential marker

b) Enhancement of tumor using antisense and triple helix anti-IGF-I technology. In this type of immunogene therapy, the tumor cells are down-regulated in production of IGF-I when transfected with vectors either expressing IGF-I antisense RNA or inducing IGF-I RNA-DNA triple helix. The transfected cells lose tumorigenicity and elicit tumor specific immunity which leads to the cure of established tumors [8].

Figure 2: Increase of CD8+11b- in PBL cells removed from blood cells of treated glioblastoma patients after every of three successive vaccinations, once a month (cytofluorometry, Becton Dickinson) [12,13].

Both IGF-I anti-gene therapies: IGF-I antisense and IGF-I triple helix, were introduced in clinical trial of glioblastoma (Cleveland, USA; Bangkok, Thailand; Bromberg, Poland-collaboration with Paris, France), liverhepatocarcinoma (Shanghai, China, and Cracow, Poland), cancers of prostate, colon, liver, uterus and ovary (NATO Science Program-USA, France, Poland, Germany), and melanoma, prostate and colon (Bogota and Bucaramanga, Colombia - clinical study in progress) [5,7-11]. The first clinical results were very promising and indicated an increase in immune response in vaccinated patients. All patients showed phenotypic changes in PBL; in general, the specific features of activation, both Th and Tc populations were observed in our results, especially an increase in CD8+ percentage with characteristics switching from CD8+CD11b+ to CD8+CD11b- phenotype. The life span of treated glioblastoma patients ranged between 18 and 24 months and in some cases the observed survival was 3 and 4 years. Similar results of CD8 typing were observed in immunogene therapy of other tumors (i.e. prostate, colon and liver cancers) [7,12] (Figure 2). In the first step of clinical trial Phase I proposed in Colombia in UNAB University/Fiscal Clinic, three to four cases of every following cancer diseases will be treated: glioblastoma, cancers of prostate and colon [12,13].

References
