Effective nivolumab sequential thoracic radiotherapy in elderly patients with advanced squamous cell lung cancer: did radiation therapy play a role? A case report

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Introduction

Non-small-cell lung cancer (NSCLC) accounts for over 80% of all lung cancers. Approximately 75% of all NSCLC cases are diagnosed at advanced stage with a median survival of 4–5 months after diagnosis and 10% of 1-year estimated survival. Furthermore, nearly 50% of all cases of lung cancer occur in the elderly subpopulation aged over 70 years, in which physiological changes together with comorbidities exert a negative impact on systemic therapy tolerability and overall outcome. Among histologies, squamous cell carcinoma represents approximately 30% of all NSCLC, the management of which, with first and second line chemotherapy, is difficult in elderly patients. A combination of two drugs in first line chemotherapy – cisplatin plus vinorelbine or weekly nab-paclitaxel with carboplatin – has shown superior efficacy over monotherapy in fit elderly patients, although this benefit occurs at the expense of the patient in terms of added toxicities. But for second line therapy little therapeutic progress has been made since the approval of docetaxel in 1999. Since then, no more effective agents have been found due to toxicity and lack of specific genetic alterations in squamous cell lung carcinoma.

Radiotherapy is acknowledged as an important partner in lung cancer treatment, but in elderly patients with advanced stage squamous cell lung carcinoma, it is not always used with curative intent. However, evidence suggests that cases with oligometastatic...
disease responding to systemic chemotherapy may be selected for curative radiotherapy.7

Advances in cancer immunotherapy and emerging knowledge of the immunogenic power of radiation might change the scenario and promote cure.

The cancerogenesis process seems to be based on altered cancer immuno-surveillance as a result of tumor cells escaping immunity,6 and there is an increased body of evidence showing that radiotherapy enhances immune responses against cancer.9 A combination of several immunotherapeutic strategies and radiation may be able to reshape the tumor’s microenvironment toward the inhibition of tumor-induced immunosuppression, leading to the immunogenic death of cancer cells.10

Promising effects are coming from therapies directed against cancer immuno-resistance mechanisms such as programmed cell death protein-1 (PD-1) signaling or its ligands (PD-L1, PD-L2) and CTLA-4.11 Among them, one important mechanism involved in tumor cells’ escape recognition and elimination by the immune system is the interaction of PD-1/PD-L1.12 These proteins are prevalently expressed in NSCLC and have been found to down-regulate T cell activation and promote tumor “immune escape”.13 Studies have demonstrated that the inhibition of the PD-L1/PD-1 pathway is effective for refractory solid tumors, but the efficiency of this treatment related to the tumor PD-1/PD-L1 expression level remains unclear and controversial.14 Experimental preliminary results suggested that PD-L1 positivity in the tumor specimen may correlate with response to treatment with anti PD-1 inhibitors.15,16 Additionally, a positive clinical outcome seems to be related to tumor-infiltrating lymphocytes (TILs) in the specimen.17 Interestingly, TILs have been found to have a prognostic and predictive value in patients with advanced NSCLC treated with platinum-based chemotheraphy.18

Current evidence indicates that radiotherapy can, via diverse cascade mechanisms, invoke both local and systemic immunoresponses promoting tumor-cell death through an in situ vaccination effect.19 These immunostimulatory properties have recently generated a widespread interest in light of the preclinical and clinical evidence of the “abscopal effect” in which localized hypofractionated radiotherapy has induced an antitumor immune response mediating the regression of non-irradiated metastases.20

Combining radiotherapy and immunotherapy could be a crucial strategy to overcome cancer immuno-resistance and improve patient survival, as we have supposed for this case report.21

Case presentation
Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review with the Editor in Chief of this journal.

In January 2015, an 80-year old Caucasian male patient came to our attention suffering from cough with blood in sputum, dyspnea, and moderate weight loss. He informed us that he had been a smoker of 20 cigarettes per day but stopped 10 years ago and he did not report alcohol consumption. At clinical examination he showed an ECOG performance status of 1, while no palpable lymph nodes in the neck and supraclavicular fossa were found. The chest X-ray recorded a mass in the right lung measuring at least 3 cm with a nodular lesion in the posterior side of the same lobe plus a nodule in the left lung’s upper lobe. The total body computed tomography (CT) confirmed a contrast enhancing irregular mass measuring 35×25 mm in the upper lobe of the right lung with the nodule in the posterior side of the same lobe near the pulmonary fissure measuring 10 mm. A nodule in the upper posterior lobe of the contralateral lung, of 27 mm, was also identified (Figure 1).

Subsequently, PET confirmed the suspicion of malignancy in both lungs: the large right lesion showed a high uptake with standardized uptake value 14.1, while the smaller one had a 2.1 SUV; the left nodule had a SUV 9.1. Brain Magnetic Resonance Imaging (MRI) scan was negative. Stage of disease was T3 N0 M1a (Stage IV) according to American Joint Committee on Cancer (AJCC) seventh edition (Figure 2).

A bronchoscopy confirmed the presence of a stenotic exophytic lesion arising from the mucosa of the sub-segmentary upper right bronchus. A biopsy on the nodule of the left lung and involved bronchus was done during bronchoscopy; the histopathologic examination identified NSCLC in both specimens.

Histopathologic examination identified a poorly differentiated (grade 3) squamous cell carcinoma associated with an infiltration of lymphocytes and macrophages distributed in the specimen. TILs were assessed only by H&E stain but CD3/CD4-CD8 subpopulations were not specified. Immunohistochemistry was employed to detect biological features such as TIF-1, CK7, and p63. Consequently, TIF-1 and CK7 negativity and p63 positivity were found.

Indirect immunoperoxidase procedures were carried out to assess the PD-1 and PD-L1 expression and, as a result, a moderate staining (2+) in more than 5% of tumor cells was observed.

Given the stage of the disease, the squamous histology, age >70 years, and performance status 1 typical of
a fit elderly patient, cisplatin-containing chemotherapy treatment scheme was prescribed. The chosen scheme consisted of cisplatin 50 mg/m$^2$ on day 1 plus intravenous vinorelbine 22.5 mg/m$^2$ on days 1 and 8 every 21 days for four cycles. Only two cycles were delivered due to acute severe toxicity – Grade 2 anemia and G3 febrile neutropenia requiring erythropoietin for 2 months and G-CSF for 3–4 days and antibiotics. After hematologic recovery, the patient continued to receive intravenous vinorelbine, as mono-chemotherapy, 25 mg/m$^2$ on days 1–8 every 3 weeks for four cycles. Acute toxicity was mild – G2 thrombocytopenia after the last cycle and constipation.
One month after the last vinorelbine delivery, restaging PET/CT was performed, only a complete remission of the two pulmonary nodules was observed, while the right intrapulmonary lesion was increased measuring 45 mm with an avid fluorodeoxyglucose (FDG) uptake with an SUV 18. The disease was downstaged to yT2a N0 M0 (stage IB) due to disappearance of the two nodules (Figure 3).

On the basis of these results, the patient was considered eligible for locoregional treatment with thoracic radiotherapy. Respiratory function tests (FEV₁ and DLCO) were required to assess the lung capacities and diffusion. Spirometry recorded a restrictive respiratory impairment with an FEV₁ 2.36 L; the alveolar-capillary diffusion was low with a DLCOc/VA 0.54 mmol/min/kPa/L (47%) due to interstitial damage. 3D conformal radiotherapy was delivered using five multileaf collimator customized isocentric fields and MV 10 photon beams. The planning target volume (PTV) consisted of the intrapulmonary post-chemotherapy right lesion as gross tumor volume (GTV) with an isotropic margin of 1.8 cm. The prescribed dose was 66 Gy with standard fractionation (2 Gy/fr). To minimize the risk of radiation pneumonitis, in the histogram dose volume plot (DVH) the constraint limit for the whole lung as the sum of both lungs was respected with V20=25%; V10=45%; V13=38%, V5=50% and mean lung dose =12 Gy. Concurrent chemoradiotherapy was excluded due to initial stage, age, and previously recorded chemotherapy-induced toxicity.

Acute radiation toxicity was moderate – G2 dyspnea occurring during the 4th week of treatment. One month after completion radiotherapy, despite the optimal lung DVH as mentioned previously, the patient developed symptomatic acute febrile grade one radiation pneumonitis. Steroids and antibiotics were effective. At second restaging 1 month after pneumonitis was resolved, a new CT scan showed only a modest reduction of the right pulmonary mass with a high contrast enhancement surrounded by interstitial fibrosis. The PET after 3 months of radiotherapy showed a metabolically active right pulmonary lesion with avid FDG uptake (SUV 14) (Figure 4). The patient was diagnosed with a refractory tumor. At that time the patient complained of moderate exertional and at-rest dyspnea, fatigue due to anemia (with Hb 10 g/dL), and weight loss of more than 5 kg. At clinical examination the patient

Figure 3 PET at first restaging after chemotherapy.
Note: The disease is downstaged with persistence of primary tumor in right lung: the stage is yT2 N0 M0.
appeared more frail, pale, and unfit for further combination chemotherapy. The stage of disease remained as yT2 N0 M0. As second line of systemic therapy, we chose to start immunotherapy on the basis of the previous PD-1 determination of the biopsy specimen. Nivolumab monotherapy 3 mg/kg intravenously every 3 weeks for a total of eight cycles was prescribed and started 4 months after radiation completion. Pruritus was the main acute side effect, no other adverse effects were recorded. During this therapy the performance status and fatigue were ameliorated with a hemoglobin level increase to 12 g/dL and a good weight gain of more than 10 kg. After six cycles of nivolumab, the patient underwent total body CT restaging which showed complete resolution of the right pulmonary lesion consisting of a tissue loss with a cavitation instead of the avidly irradiated mass surrounded by an extended pulmonary fibrosis in the medio-apical part of the right lung.

After eight cycles the new PET confirmed the complete resolution of the tumor, as well as no FDG uptake (Figure 5). After 24 months (the last nivolumab delivery), the PET and CT were still negative for local or systemic disease recurrence. The patient, who is now 83 years of age, has had no disease recurrence (as of December 2017), with a good PS, and has regained quality of life.

Discussion

NSCLC is the most common cancer in the world and the leading cause of cancer-related deaths in Western countries. Approximately one third of lung cancers are diagnosed in patients older than 75 years and the majority of them already have metastatic disease. Palliation is the main or the only therapeutic option. In regard to an effective systemic approach to this population, some Phase II randomized trials confirmed the cisplatin combination plus vinorelbine as a feasible and active treatment in elderly patients with advanced NSCLC. Vinorelbine alone or with cisplatin for physically fit, chemo-naïve NSCLC patients older than 70 years, has been defined effective with a better response rate and a longer median time to disease progression. However, both statistically significantly higher toxicity and no survival advantage for the combination treatment have been observed.

Our patient was treated according to this currently available evidence-based therapy and developed acute hematologic toxicity as expected, and a poor outcome. The only evaluable positive effect was downstaged disease which consisted of the resolution of the pulmonary nodules, which meant he was eligible for thoracic radiotherapy. However, radiotherapy did not yield a curative result. This scenario and the conclusions to be drawn could be discouraging because an elderly patient with advanced squamous lung cancer, whose disease does not respond to first line chemotherapy and curative radiotherapy, has limited chances of cure. Finally, the proven existence of altered immunosurveillance in cancer development and the availability of immunotherapy could reshape this scenario, as we have hypothesized in this case.

An old hypothesis of active immunosurveillance that protects against cancer has been revalued and most recently refined into the process termed “cancer immunoediting”, in which tumor cells and immune cells interact within its microenvironment through three E phases (elimination, equilibrium, escape) and by prevalence of one of them, the tumor’s fate depends. Research has revealed the ability of tumors to escape the immune system through the activation of distinct pathways including endogenous “immune check points” such as targeted proteins, eg, PD-1, PD-L1, and CTLA-4, which exert a negative regulation on immune anticancer response, permitting cancer progression and metastasis. Advances in immunotherapy have resulted in therapies against these immune check points, the so called immune check point inhibitors, which have achieved promising results in refractory metastatic solid tumor-affected patients.
Nivolumab, pembrolizumab, ipilimumab, tremelimumab, and durvalumab, at last, are all immune check point inhibitors capable of binding and inactivating the effects of the previously mentioned proteins, leading to immunogenic cancer cell death, with a safe toxicity profile.\(^{30,31}\) Moreover, preliminary reports suggest that factors such as PD-L1 expression, presence of TILs, and smoking, correlate with response to these therapies.\(^{32–34}\) All these factors were present in our patient.

Nivolumab is a fully humanized IgG4 monoclonal antibody (mAb) that targets PD-1, and its successful use as a second line therapy in NSCLC has led to its definitive US Food and Drug Administration (FDA) approval as second line treatment for treatment-resistant patients,\(^{35,36}\) such as our patient. Despite the encouraging results, the response rate to immunotherapies is still low; in fact, positive outcomes have been recorded in approximately 17% to 19% of unselected patients, probably due to low immunogenic tumor expression in such cases.\(^{37,38}\)

This has led to the search for strategies to overcome this resistance, modifying the tumor immune microenvironment. Data from many laboratories and sporadic reports indicate that local radiation therapy exerts immunostimulatory properties resulting in systemic, immune-mediated antitumor and distant anti-metastatic effects (abscopal effect) through increased tumor-antigen release and antigen-presenting cell cross-presentation, improved dendritic cell function, and enhanced T-cell priming.\(^{19}\) Preclinical and clinical reports have speculated regarding the concept of radiation-induced in situ vaccination as a result of radiation-induced expanded release of tumor-associated antigens and an amplified chemokine cascade (DAMPs), which causes the immunogenic death of cancer cells.\(^{39,40}\) As a result of mouse tumor models and clinical reports of more frequent abscopal responses in patients refractory to immunotherapy who received large sized hypofractionated RT during immunotherapy, the ability of RT to enhance antitumor T cell activity in synergy with immune check point inhibitors has been acknowledged.\(^{41}\) Moreover, some authors have suggested that, in patients who would otherwise not respond to

**Figure 5** PET at last restaging after eight cycles of nivolumab.

**Note:** No fluorodeoxyglucose uptake and cavity with fibrosis in right lung is present, confirming a complete remission of disease.
immune check point inhibitors, localized radiotherapy can induce activated tumor-specific T cells, thereby promoting response to these therapies.62 Anti-PD1 therapies are generally well-tolerated in clinical trials, however concerns exist about increased adverse events with radiotherapy and immunotherapy combinations, like high grade pneumonitis among NSCLC patients who previously received thoracic radiotherapy. The incidence of high grade pneumonitis in this patient population has been estimated as 4.1%, suggesting the need for closer toxicity monitoring among patients who were previously treated with chest radiation.31 Interestingly, our patient developed pneumonitis after the completion of radiotherapy, before nivolumab treatment started. These arguments suggest a role of radiotherapy-induced inflammatory activity in priming and homing of activated immune cells in the irradiated area leading to a favorable microenvironment more sensitive to the sequential anti-PD1 activity. In support of this hypothesis, experimental data have shown how fractionated radiotherapy leads to an adaptive up-regulation of tumor cell PD-L1 expression that is dependent on CD8 T cells’ production of IFNγ. This effect seems to attenuate the anticancer radiation immune response that can be overcome by the combination with mAB PD1-PDL1.44 Furthermore, Adams et al found that approximately 30% of irradiated patients with NSCLC had a substantial increase of PD-L1 expression in circulating tumor cells 14–21 days after radiotherapy. In a sequential analysis of patients 2–4 months after completion of radiotherapy, nearly 87% of the PD-L1 expression remained unchanged from the T1 time point.45 These findings could be the rationale underlying the effect of nivolumab given 4 months after radiotherapy in our patient.

Going from the experimental to clinical data, currently, over a dozen clinical trials have evaluated the activity of anti-PD1 antibodies in previously irradiated refractory NSCLC advanced stage patients. Checkmate 017 trial enrolling stage III–IV NSCLC pretreated patients reported that, compared to docetaxel, nivolumab achieved superior overall survival.46,47 It is curious that in this trial there was no correlation between PD-L1 expression and the benefit of nivolumab, and only one patient in the nivolumab arm (out of 272 enrolled patients) achieved a complete response. Nevertheless, from the study design it is not clear whether previously irradiated patients were included, and up to now no subgroup analysis data are available yet. An interesting suggestion regarding this concern has been put forward by the secondary analysis of the KEYNOTE-001 Phase I trial involving the group of previously irradiated NSCLC patients receiving pembrolizumab, which is another mAb that targets PD-1.48 In this analysis, patients receiving any radiotherapy to any site achieved a better and prolonged median progression-free survival and overall survival than non-irradiated patients. On the other hand, more recently the Pacific trial has reported exciting results with durvalumab vs placebo in a homogeneous population all pretreated with chemoradiation.49 Therefore, there are substantial data to support the hypothesis of the synergistic role of radiotherapy and sequential nivolumab, as shown in our patient in whom PD-L1 expression was positive but low. If so, the underlying mechanism could be as assumed by Demaria et al, who explained how radiotherapy is able to convert an immunogenic “cold” tumor into an immunogenic “hot” tumor, changing the game of immunotherapy.50 Certainly factors such as smoking, PD-L1 expression, and TILs have given an effective contribution to achieve this result.

Conclusion

Elderly patients with advanced NSCLC whose disease progresses or does not respond to first line chemotherapy and/or local radiotherapy have limited therapeutic options. The introduction of sequential or concomitant immunotherapy with conventional radiotherapy could be an effective therapeutic option to overcome cancer immune resistance and improve cancer-specific survival as emphasized by experimental models,44 Phase I–II clinical trials47–48 and sporadic reports, and more recently by a Phase III clinical trial.49

Apart from clinical trials, this case report confirms the experimental and preliminary data on the synergistic effects of nivolumab and radiotherapy in a heavily pretreated squamous cell lung cancer patient who had a poor response to radiotherapy. Robust Phase II–III clinical studies are required to better support these arguments. In fact, a Phase II safety-efficiency European Clinical Trial and Phase III RTOG 3505 trial providing nivolumab after standard chemoradiotherapy in stage III NSCLC patients have been designed, although, up until now they have not been open for enrollment.51,52

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Author contributions

All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.
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The authors report no conflicts of interest in this work.

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