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**EFGR-mutant lung adenocarcinoma and Li-Fraumeni syndrome: report of two cases and review of the literature.**

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Abstract:

We report two cases of non-smoker patients diagnosed with EGFR-mutated lung adenocarcinoma and bearing germinal TP53 gene mutation, also known as Li-Fraumeni syndrome (LFS). We describe for the first time an EGFR-TKI resistance mutation in this population. Finally, we provide an analysis of discerning epidemiological data obtained from the IARC database and from all the published cases of EGFR-mutated lung cancer in TP53 germline mutation carriers.
Introduction

Li-Fraumeni syndrome (LFS) was initially described in the late 60’s, based on the report of four families with high tumor occurrence [1]. This condition has been characterized by an autosomal dominant predisposition to various malignancies. The principal localizations are sarcomas, gliomas and breast carcinomas. Direct involvement of germline TP53 gene mutations has emerged approximately two decades after the primary description [2]. In most cases, mutation of the TP53 tumor suppressor gene result in a loss-of-function of p53 protein, leading to alterations in DNA repair, apoptosis cell-cycle control and consequently genome instability [3]. A worldwide database is now collecting both somatic and germline TP53 mutation on International Agency for Research on Cancer (p53.iarc.fr). Despite growing knowledge about LFS, prevalence of lung cancer is still unknown in this population but an increased risk has been described in observational studies [4]. Moreover, the literature is still scarce regarding the molecular abnormalities of lung neoplasms in this particular setting. An early report suggests an association between tumoral TP53 mutation and EGFR-driven lung cancer [5].

We report two cases of patients with LFS and EGFR-mutated lung adenocarcinoma, and provide the first description of resistance mutation to EGFR-TKI in this population. We highlight a potentially underreported association and emphasize the need for molecular alterations compiling in the particular setting of LFS. Reviewing all the published cases to date, we finally bring perspectives for clinical management.
Cases

A fifty-five year old non-smoker man was diagnosed with a lower left leg deep venous thrombosis in December 2010. CT-scan evaluation revealed upper right lobe lung mass, abnormal mediastinal lymph nodes and two liver lesions. A liver biopsy confirmed the presence of a metastatic localization of a stage IV (T2a N3 M1b) lung adenocarcinoma. Eastern cooperative oncology group (ECOG) performance status was quantified to 0 at diagnosis. The patient was known to carry a LFS, because emergence of unusual childhood tumors in descendants led to germline TP53 gene sequencing a few years ago (TP53 germline mutation H179Y). First line chemotherapy with platinum and pemetrexed was administered, followed by EGFR-TKI (erlotinib). Three months later, paclitaxel was prescribed because of a cervical and mediastinal progression, which stabilized the disease for eight months. Symptomatic pleural effusion occurred and the patient clinically benefit from a re-challenge of platinum doublet chemotherapy. Approximately two years after diagnosis, CT-scan guided biopsy was performed on a new pulmonary nodule to assess tumoral molecular abnormality status and rule out transition to small cell lung cancer. The presence of an adenocarcinoma with both an \( EGFR \) exon 19 deletion (del2235–2249, delE745-A750) and an exon 20 T790M (c.2369C>T) mutation was confirmed respectively by automated electrophoresis system (LabChip GX) and pyrosequencing (figure 1, panel A and B). A tumoral \( TP53 \) gene mutation was also found, similarly to the germline mutation previously described (data not shown). No mutation on \( HER2, PI3KCA, BRAF \) or \( KRAS \) genes or rearrangement of ALK gene was found. A fourth line of chemotherapy was proposed with an objective response observed after 4 cycles of carboplatin pemetrexed bevacizumab. The patient died a few months later from rapidly progressive pulmonary lymhangitic carcinomatosis (figure 1, panel C).
A fifty-seven year old woman presented with a two month history of inflammatory lumbar pain in November 2008. MRI evaluation revealed a body lesion of L1 suspect of malignant origin. Needle bone biopsy confirmed the presence of a secondary metastasis of an upper left lobe lung adenocarcinoma. It was a stage IV (T2aN2M1b) disease and ECOG performance status was quantified to 0 at diagnosis. She never smoked and presented right breast carcinoma in 2002, treated by surgery and radiation. EGFR status was unknown at diagnosis because bone sample conditioning does not allow us to perform molecular analysis. After two cycles of cisplatin vinorelbin, the disease progressed, therefore erlotinib was prescribed. This therapy resulted in a significant and prolonged response. In October 2009, she was tested for germline TP53 mutation because her daughter was diagnosed with corticosurrenaloma. LFS was confirmed (TP53 mutation R273H). Thoracic progression occurred in November 2011 and the patient was enrolled in a phase 1 study evaluating an irreversible EGFR-TKI inhibitor (afatinib). Pleural metastasis was diagnosed in March 2012 and the targeted therapy was stopped. A surgical pleurodesis showed pleural localization of an adenocarcinoma with EGFR exon 21 mutation L858R (c.2573T>G) without T790M mutation, as assessed by pyrosequencing (figure 1, panel D). No mutation on HER2, PI3KCA, BRAF or KRAS genes or rearrangement of ALK gene was found. Consequently, a re-challenge with erlotinib resulted in a 5 month stable disease. The patient died from respiratory failure because of multiple pulmonary metastasis, more than four years after initial diagnosis (figure 1, panel E).
Discussion

We report two LFS patients diagnosed with EGFR-mutated lung adenocarcinoma. Observational studies show that lung cancer belong to the spectrum of predisposed tumors in LFS [4]. According to Chompret criteria revised in 2009, a proband diagnosed with lung cancer before the age of 46 years and with at least one first-degree or second-degree relative with a core LFS tumor should be selected for a TP53 germline mutation testing. Our two patients do not meet these criteria and LFS was evoked because early-onset cancers in family history. Approximately 29%–35% of families with these criteria were shown to have TP53 germline mutations [6][7]. Importantly, recognition of LFS impacts the familial and personal cancer risk assessment, regarding potential radio-induced neoplasms. Thoracic oncologists should be aware of the possibility of this inherited predisposition syndrome especially when familial history of early-onset cancers is known. Our cases highlight the need for exhaustive medical family history interrogation when lung cancer is diagnosed, notably for young and non-smoker patients.

The literature is scarce regarding epidemiologic characteristics of lung cancer in LFS. The International Agency for Research on Cancer mutation database collecting the germline TP53 mutation reports various pathological subtypes in this restricted population [8]. Adenocarcinomas, including lepidic growth carcinomas (previously known as bronchoalveolar), represent the largest part of the confirmed pathological type when TP53 germline mutation is known (figure 2, panel A). In addition, majority of these tumors arise before the fifth decade and for a large part in women (figure 2, panel B). To date, this database do not contain molecular characteristic of these tumors. In our two patients, EGFR mutational status was unknown at diagnosis, but found during follow-up. Interestingly, co-
occurrence of tumoral somatic \textit{TP53} mutations and \textit{EGFR} mutations has been found in 34\% of lung adenocarcinomas in a recent international project [9]. Two retrospective studies tried specifically to assess this association, based on mixed genomic and immunohistochemical approaches [5][10]. Tumoral p53/arf pathway inactivation was observed in respectively 100\% and 58.1\% of \textit{EGFR}-mutated lung cancers, supporting this hypothesis. Interestingly, a similar distribution of germinal and tumoral mutations is observed in the \textit{TP35} gene coding sequence, occurring preferentially in the DNA binding domain (exon 5 to 8). Nevertheless, it appears that all tumoral \textit{TP53} mutations are not equivalent regarding their intrinsic prognostic value. Indeed, a recent publication has described that non-disruptive \textit{TP53} mutations are associated with a worse outcome in \textit{EGFR}-mutated NSCLC [11]. Moreover, a recent analysis of age of cancer onset from the IARC database suggests unequal repercussion of the different \textit{TP53} germline missense mutations [12]. Taken together, these clinical data suggest an important heterogeneity concerning the tumorigenic potential of the different subtype of \textit{TP53} mutations.

No exhaustive data concerning molecular biologic characteristics of lung cancers in the context of LFS has been published so far. Nevertheless, a growing number of publications reported \textit{EGFR} mutant lung cancer in carriers of LFS suggesting a potential link between these two conditions [13][14][15][16]. Table 1 recapitulates important characteristics of the previous cases reported as well as our two patients. Consistent with IARC database, the majorities of tumors arise before the fifth decade and occur predominantly in woman. Interestingly, a majority of LFS patients harboring \textit{EGFR} and/or \textit{ERBB2} mutated lung adenocarcinomas encompass good tumoral and clinical response to targeted therapy (except for the patient with T790M resistance mutation). The benefit appears to be quite similar than in non-LFS population with a progression-free survival of approximately one year. Some dedicated studies are needed to confirm this observation. As expected from clinical trials,
rechallenge with EGFR-TKI after progression show no benefit in our second case, even in the absence of identified resistance mutation. Third generation TKI recently show promising anti-tumor activity, but current investigation are limited to the tumors with identified resistance mutation. Furthermore, we report for the first time acquisition of EGFR-TKI resistance mutation in the context of LFS. Familial cases of germline T790M may exist in 50% of these cases [17]. It was ruled out in our first patient because allele frequency was not consistent with this hypothesis (around 7%) and the mutation was not found in her daughter genome (data not shown). However, we cannot exclude that EGFR T790M mutation was not present at the time of diagnosis because reduced sample materials from hepatic needle-biopsy do not allow us to perform molecular biology. Still, these cases suggest a common natural history of resistance to TKI than in general population. In LFS patients, as the first genetic hit is inherited, we argue that a second-hit, according Knudsen hypothesis, can arise sooner in the time course of tumorigenesis. Interestingly, some authors incriminate radiation therapy or chemotherapy as a second hit. Notably, five of six LFS cases with EGFR-mutated lung adenocarcinoma were previously treated with chemotherapy or radiotherapy in a context of mammary malignancies. Even if a causative role remains to be demonstrated, benefice-risk assessment of radiotherapy should be carefully evaluated and discussed with patients in this particular setting.

In conclusion, thoracic oncologists should evoke LFS in the context of illegitimate lung cancer (young and non-smoker patients). EGFR mutation is frequently reported in the Li-Fraumeni setting, but epidemiological studies are needed to assess this potential association. Response to targeted therapy seems to be similar as in the general population. Arguably, we show for the first time that same resistance mechanism could arise in the LFS context.
References


Author contributions:

CR, MLT, MK, HL and BD were responsible for the literature search. CR, MLT, MK and HL were responsible for figures. CR was responsible for data collection. CD, AL and JM were responsible for molecular biology data collection and analysis. All authors were responsible for data interpretation, and writing and final approval of the article. CR takes responsibility for the paper as a whole.

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Legends:

Figure 1:

(A) EGFR exon 19 deletion diagnosed by automated electrophoresis system (LabChip GX, Perkin Elmer®). (B) EGFR T790M resistance mutation (ACG→ATG) as assessed by pyrosequencing of tumoral DNA. (D) EGFR L858R activating mutation (CTG→CGG) as assessed by pyrosequencing of tumoral DNA (*: indicate mutated nucleotide or deleted allele). (C, E) Evolution of the sum of the longest diameters for all targeted lesions based on CT-scan measurement during follow-up.

Figure 2:

(A) Pie chart of pathological types of lung neoplasm in germline TP53 mutation carriers (n=34) (†: including the bronchioloalveolar carcinoma). (B) Diagram of age and sex repartition of lung cancer in LFS patients (n=33; age missing for one patient). Data extracted from the International Agency for Research on Cancer p53 mutation database [9].

Table 1:

<table>
<thead>
<tr>
<th>Age</th>
<th>Gender</th>
<th>Smoking status</th>
<th>Medical histories</th>
<th>Histology</th>
<th>Stage</th>
<th>TP53 mutation</th>
<th>Driver mutation</th>
<th>Response to targeted therapy</th>
<th>Resistance mechanism</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>55</td>
<td>M</td>
<td>Non-smoker</td>
<td>- None</td>
<td>Poorly differentiated invasive right lung adenocarcinoma</td>
<td>IV</td>
<td>Mutation H179Y</td>
<td>EGFR deletion in exon 19</td>
<td>PD</td>
<td>EGFR mutation T790M</td>
<td>This work</td>
</tr>
<tr>
<td>57</td>
<td>F</td>
<td>Non-smoker</td>
<td>- High grade ductal right breast carcinoma treated by tumorectomy and radiation</td>
<td>Moderately differentiated invasive left lung adenocarcinoma</td>
<td>IV</td>
<td>Mutation R273H</td>
<td>Mutation EGFR L858R</td>
<td>PR (one year - erlotinib)</td>
<td>-</td>
<td>This work</td>
</tr>
<tr>
<td>34</td>
<td>F</td>
<td>Non-smoker</td>
<td>- Extensive ductal right breast carcinoma treated with surgery, adjuvant chemotherapy, autologous stem-cell transplantation, and radiation</td>
<td>Invasive left lung adenocarcinoma (with broncho-alveolar feature)</td>
<td>IV</td>
<td>Mutation R273H</td>
<td>EGFR deletion in exon 19</td>
<td>Not specified (one year - erlotinib)</td>
<td>-</td>
<td>[13]</td>
</tr>
<tr>
<td>37</td>
<td>F</td>
<td>Non-smoker</td>
<td>- Malignant phyllodes tumor and bilateral ductal breast carcinoma in situ treated by bilateral mastectomy and adjuvant tamoxifen</td>
<td>- 2008: moderately differentiated invasive left lung adenocarcinoma (with broncho-alveolar feature)</td>
<td>IV</td>
<td>Mutation R248W</td>
<td>EGFR exon 20 insertion (A767_S768)</td>
<td>PR (6 months - lapatinib with decotaxel and lapatinib with trastuzumab)</td>
<td>-</td>
<td>[14]</td>
</tr>
<tr>
<td>46</td>
<td>F</td>
<td>Non-smoker</td>
<td>- Suspected tuberculosis infection - Invasive ductal breast carcinoma treated by chemotherapy - Gluteal schwannoma</td>
<td>Moderately differentiated right lung adenocarcinoma</td>
<td>IV</td>
<td>Mutation G245S</td>
<td>Mutation EGFR L858R</td>
<td>CR (one year - afatinib)</td>
<td>-</td>
<td>[15]</td>
</tr>
<tr>
<td>51</td>
<td>F</td>
<td>Non-smoker</td>
<td>- Multiple right breast invasive ductal carcinoma treated with radiotherapy, mastectomy, then chemotherapy - Right flank lump malignant fibrous histiocytoma. - Left breast invasive ductal carcinoma treated by mastectomy and chemotherapy</td>
<td>Poorly differentiated invasive right lung adenocarcinoma</td>
<td>IV</td>
<td>Deletion in exon 5</td>
<td>Mutation EGFR L858R</td>
<td>PR (one year - erlotinib)</td>
<td>-</td>
<td>[16]</td>
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