

Early occurrence of lung adenocarcinoma and breast cancer after radiotherapy of a chest wall sarcoma in a patient with a de novo germline mutation in *TP53*

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Abstract We report a 26-year-old female patient who was diagnosed within 4 years with chest sarcoma, lung adenocarcinoma, and breast cancer. While her family history was unremarkable, DNA sequencing of *TP53* revealed a germline de novo non-sense mutation in exon 6 p.Arg213X. One year later, she further developed a contralateral ductal carcinoma in situ, and 18 months later a jaw osteosarcoma. This case illustrates the therapeutic pitfalls in the care of a young cancer patient with *TP53* de novo germline mutations and the complications related to her first-line therapy. Suggestion is made to use the less stringent Chompret criteria for germline *TP53* mutation screening. Our observation underlines the possibly negative effect of radiotherapy in generating second tumors in patients with a *TP53* mutation. We also present a review of six previously reported cases, comparing their cancer phenotypes with those generally produced by *TP53* mutations.

Keywords Li-Fraumeni syndrome · Germline mutation *TP53* · De novo mutation · p.Arg213X

Introduction

Li-Fraumeni syndrome (LFS) is one of the most severe forms of familial cancer susceptibility syndrome, characterized by an early age of tumor onset and a wide tumor spectrum. According to its first description [1], diagnosis should be suspected in the presence of a bone or soft tissue sarcoma in a patient under 45 years of age and a family history with a first-degree relative with any cancer under 45 years, or a sarcoma at any age. However, LFS is rarely diagnosed in routine clinical practice, and the French LFS working group has elaborated practical criteria (Chompret criteria, Table 1), which were recently updated [2]. These criteria are less stringent than the original LFS criteria [1] and have a higher sensitivity that could allow more cases to be detected.

No other genes than *TP53* have been convincingly associated with LFS [3]. Using current techniques (sequence analysis, deletion testing), *TP53* mutation is detected in about 70% of LFS families [4]. Molecular diagnosis of these cases is critically important to ensure an appropriate medical follow-up and to avoid inappropriate cancer treatment such as radiation therapy. De novo mutations of *TP53* represent a clinical pitfall, since the mutational status is not suspected at the time of first cancer diagnosis.

We report here a case of a woman harboring a de novo *TP53* mutation, diagnosed with a chest wall sarcoma, who subsequently developed lung and breast cancer, one of them in the radiation fields, and later contralateral ductal carcinoma in situ (DCIS) and jaw osteosarcoma.

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Table 1 2009 Chompret criteria for germline *TP53* mutation screening

- I. Proband with tumor belonging to LFS tumor spectrum (e.g., soft tissue sarcoma, osteosarcoma, brain tumor, premenopausal breast cancer, adrenocortical carcinoma, leukemia, lung bronchoalveolar cancer) before age 46 years AND at least one first- or second-degree relative with LFS tumor (except breast cancer if proband has breast cancer) before age 56 years or with multiple tumors; OR
- II. Proband with multiple tumors (except multiple breast tumors), two of which belong to LFS tumor spectrum and first of which occurred before age 46 years; OR
- III. Patient with adrenocortical carcinoma or choroid plexus tumor, irrespective of family history

LFS Li-Fraumeni syndrome

Report

A 29-year-old woman with a personal history of chest wall sarcoma and second primary lung adenocarcinoma was referred to the oncogenetic clinic for a suspicion of LFS. Three years earlier, she presented a right chest wall lesion beneath the right breast. Biopsy of the lesion revealed a low-grade spindle cell sarcoma. She was treated with a wide surgical excision, axillary lymph node dissection, and adjuvant radiation to the right chest wall. She received 50 Gy in 25 fractions over 37 days. The radiation field

included the outer quadrants of the right breast that received 80% of the radiation dose (Fig. 1).

Twenty-seven months later, a follow-up thoracic and abdominal CT-scan revealed a pulmonary nodule in the superior segment of the left inferior lobe (Fig. 1). The patient underwent a diagnostic wedge resection of the pulmonary nodule, and pathology revealed a 4-mm bronchioloalveolar carcinoma. Of note, the patient had never smoked.

Family history (Fig. 2) disclosed a maternal aunt with a choledochal cancer at age 52 and a sister of her grandmother affected with colon cancer at age 79. The classical diagnostic criteria of a LFS were not fulfilled, but the occurrence of a sarcoma and lung bronchioloalveolar cancer before age 46 years met the updated Chompret criteria, prompting molecular genetic screening for a *TP53* germline mutation. After obtaining the patient's written informed consent, a peripheral blood sample was taken to perform analysis of the *TP53* gene from genomic DNA. Following PCR amplification and bidirectional sequencing of all coding exons (including intron/exon junctions) of the *TP53* gene (Ensembl-ID: ENSG00000141510), a non-sense c.637C > T (p.Arg213X) mutation in exon 6 was identified. Sequencing of exon 6 in her parents and her brother showed no such variant, indicative of a de novo mutation in the patient.

Thirty-nine months after the initial sarcoma diagnosis and at the same time of the molecular genetic confirmation

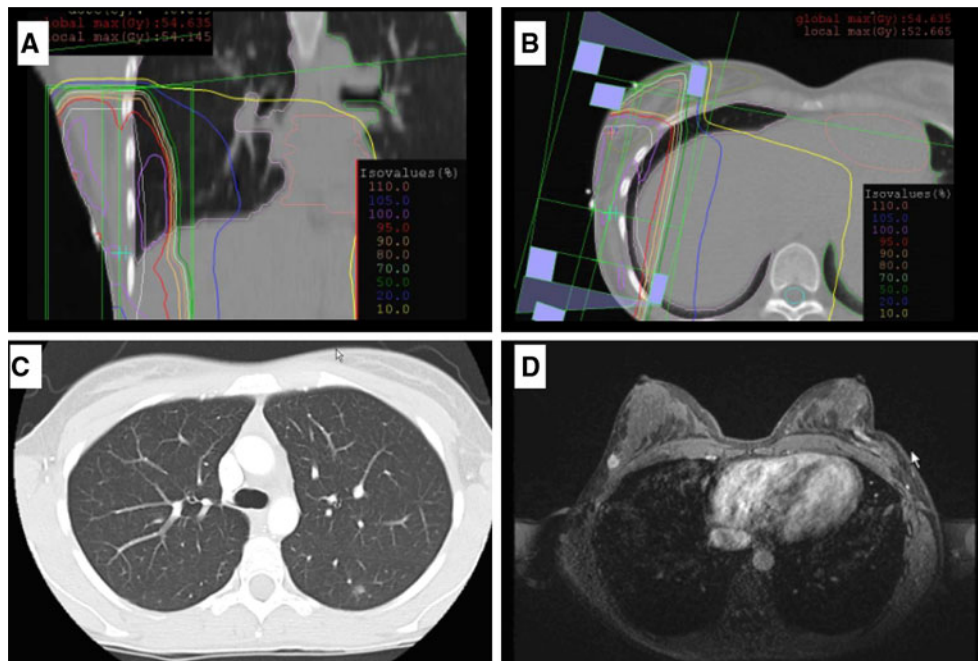


Fig. 1 **a** Isodose at level of the subsequent lung cancer. **b** Isodose at level of the subsequent breast cancer. **c** Contrast enhanced axial chest CT image showing a pulmonary nodule in the superior region of the

left inferior lobe. **d** Contrast enhanced (Gadolinium) axial T1-weighted MRI showing the tumor in the *upper-outer* quadrant of the right breast

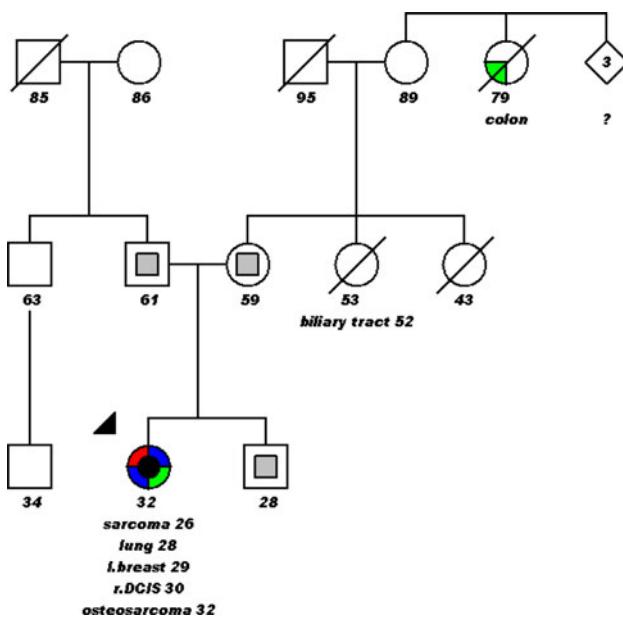


Fig. 2 Pedigree of the p.Arg213X *TP53* family: The numbers indicate the current age of the patient and the age at cancer diagnosis, the gray squares indicate the wild-type status, the black circle indicates the mutated status; DCIS ductal carcinoma in situ

of the LFS, the patient presented an upper-outer quadrant lump of her right breast on clinical examination (Fig. 1). Breast magnetic resonance imaging showed a 2-cm Gadolinium-enhanced lesion, and a biopsy confirmed a ductal carcinoma of the breast. Mastectomy of the right breast with sentinel lymph node biopsy was performed. The pathology disclosed a 1.8 cm in size, poorly differentiated infiltrating ductal carcinoma, with a negative staining for estrogen and progesterone receptors, but a positive staining for the epidermal growth factor receptor 2. The staging work-up (chest and abdomen CT-scan, bone scan, and liver MRI) was negative. A combined adjuvant treatment with docetaxel, carboplatin, and trastuzumab was initiated. However, the patient refused to continue her treatment after completion of two cycles.

One year later, during the follow-up evaluation, breast MRI showed a 10-mm contrast enhancement in the upper-outer quadrant of the left breast. The patient refused a biopsy and requested a mastectomy, which was performed with sentinel nodes biopsy. The pathological report confirmed a 35-mm large DCIS. Sentinel nodes were negative. No additional treatment was proposed. Eighteen months later, jaw pain led to the diagnosis of an osteoblastic bone lesion. A PET-scan (positron emission tomography) did not reveal metastases but strengthened the suspicion of a new primary cancer. Histological analysis of the lesion disclosed an intermediate grade 2 differentiated osteosarcoma. The patient began neoadjuvant chemotherapy. She is now 32 years old.

Discussion

Using the less stringent Chompret criteria, we have identified a patient harboring a *TP53* de novo mutation. In the initial French cohort, a *TP53* mutation was identified in only 17% of the families, using the first Chompret criteria [5]. In contrast, in a recent Spanish cohort, 95% of patients (71/75) with a *TP53* mutation met the new Chompret criteria [6]. Clinical utility of the new criteria has been confirmed by many authors who also reported *TP53* mutations in young patients with malignancies typical for LFS, but without a pathognomonic family history fulfilling the LFS criteria [7–9]. For example, in a study of 14 cases of childhood adrenocortical tumors, most germline *TP53* mutations carriers did not have a family history fulfilling the classic LFS criteria, leading to the hypothesis of a de novo mutation or a low penetrance phenotype [10]. Our patient is another perfect example where younger patients (<46 years of age) afflicted with multiple primary cancers should be strongly considered for *TP53* genetic testing even without a positive family history.

Given the very aggressive phenotype observed in our patient, it is unclear if the p.Arg213X *TP53* mutation could be responsible for that particular phenotype. The p.Arg213X *TP53* mutation is a non-sense mutation located in exon 6 and has been reported in 287 tumors. Most of the reported somatic and germline *TP53* mutations are missense mutations; only 6% are represented by non-sense mutations. The translated p53 protein, if not subjected to non-sense-mediated decay, is truncated and has lost most of the DNA-binding domains that are located downstream of amino acid 213. In the IARC *TP53* mutations database (<http://www-p53.iarc.fr/p53main.html> database version R14, November 2009) [11], the p.Arg213X mutation has been reported in 6 families [3, 9, 10, 12–17]. Five of the six articles report clinical and family data in sufficient details and are summarized in Table 2. In one report, the exact relative degree is not mentioned [3]. Finally, in all reported families, genetic testing of relatives is not reported.

Trahair et al. recently reported a female patient who had DCIS at age 30 and 34, low-grade astrocytoma at age 36, and spindle cell sarcoma at age 36 [17]. This case report shares many similarities with ours. These two patients are both female, young at first cancer diagnosis (26 and 30 years, respectively), and both had several primary tumors. The case reported by Trahair is a not a de novo mutation, since many family members presented typical LFS cancers.

The usual phenotype–genotype correlation studies have focused on *TP53* mutation presence, location, or type of mutation. An IARC database study of families fulfilling the LFS criteria demonstrates that age at breast cancer diagnosis is younger if a *TP53* mutation is present. Brain

Table 2 Cases with a p.Arg213X mutation as reported in the IARC *TP53* mutations database

Reference	Type of cancer (index case) (age at diagnosis (years))	Family history (cancer type (relative age))
Frebourg [15]	Breast (27)	Colon (N 9)
Reifenberger [16]	Cerebral PNET (<1)	Cerebral PNET (S 5), ovary (M ?), brain (U 14, U 30), other (colon, leukemia)
Varley [10]	Adrenocortical (<2)	Brain (GM 42)
Vahteristo [16]	Breast (28)	Sarcoma (64), melanoma (67), lung (75) ^a
Wong [9]	Colon (9)	Gastric (M 29), sarcoma (GFm 17), breast (Am 27)
Trahair [17]	Rhabdomyosarcoma (2)	Breast and other cancers ^b (M 30, 34), fibrosarcoma (GFm 43), ACC (GCm 8), esophageal (GUm 40), liver (GGFm 20)
Our case	Sarcoma (26) Lung (28) Breast (29) Contralateral DCIS (30) Osteosarcoma (32)	

A aunt, ACC adrenocortical cancer, C cousin, DCIS ductal carcinoma in situ, G grand, m maternal, M mother, N niece, p paternal, PNET primitive neuroectodermal tumor, S sibling, U uncle

^a Relatives degree were not reported

^b Mother had DCIS at age 30 and 34, low-grade astrocytoma at age 36, spindle cell sarcoma at age 36

tumors are associated with missense *TP53* mutations located in the DNA-binding loop that are in contact with the minor groove of DNA; adrenal gland carcinomas with missense mutations located in the loops opposing the protein–DNA contact surface and null phenotype (absence of the protein or loss of function) with earlier onset brain tumors [18]. The most significant defect is loss of function because large deletions encompassing the whole *TP53* gene have been found in LFS families with aggressive features [19]. The p.Arg213X *TP53* mutation causes the loss of most of the DNA-binding domains that are located downstream of amino acid 213, which could explain the severe phenotype of our case.

Besides *TP53* mutations, other genes can act as genetic modifiers. Many other genetic modifiers or other mechanism, not yet available for clinical practice, have been recently identified in LFS families. Polymorphic genetic variants in the p53 stress response pathway could determine an individual's susceptibility to cancer. In the p53 pathway, the *MDM2* gene encodes an important negative regulator of p53. A single nucleotide polymorphism in the promoter region of *MDM2* (SNP309) leads to a functional attenuation of p53. In 88 LFS patients harboring a pathogenic *TP53* mutation, presence of SNP308 was associated with an earlier tumor onset [20].

DNA rearrangements, such as DNA copy number variation (CNVs), can also predispose cancer development when they encompass, overlap, or disrupt dosage-sensitive genes. *TP53*, as guardian of the genome, is involved in the very processes known to give rise to CNVs, such as suppression of homologous recombination [21]. Genomic

CNVs analysis in a LFS cohort using high-density oligonucleotide arrays has shown that *TP53* mutation carriers displayed a significant increase in CNVs than in *TP53* wild-type LFS patients [22].

Among the factors described above, the loss of the DNA-binding domains located downstream of amino acid 213 explains the high number of primary cancers in our patient. To the best of our knowledge, no specific *TP53* mutation or modifying genes have explained this “multiple cancers” phenotype. We are aware of only one case report which showed the existence of a double pathogenic mutation of *TP53* and *BRCA2* [23]. The patient was identified through systematic *BRCA1*, *BRCA2*, *TP53*, *CDKN2A* germline mutation screening of 82 patients diagnosed with breast cancer and melanoma. A coexisting *TP53* mutation and *BRCA2* germline mutation was identified. This patient had developed 6 different primary cancers by age 70.

Finally, we have to strongly consider the potential deleterious influence of the radiotherapy in the rapid development of second primary cancers. High sensitivity to radiation-induced cancers has been reported *in vitro* [24] and in many *TP53* mutation carriers series [25, 26]. In the largest series of 200 Li–Fraumeni syndrome family members diagnosed with cancer, 30 had multiple primary cancers and 8 cases occurred in radiation fields at 3 to 22 years after the treatment [27]. *TP53* is involved in various cellular processes such as the regulation of DNA repair and programmed cell death. Our patient presented a lung cancer close to the radiation fields 27 months after radiotherapy, a right breast cancer in the irradiated field 39 months

after radiotherapy, as well as a left DCIS also close to the radiation fields 53 months after radiotherapy. This exposure to radiation therapy probably explains the short time elapsed between the four primary cancers in our patient. The jaw osteosarcoma, the fifth primary cancer, is the only one to be located outside of the chest and occurred 6 years after the chest sarcoma. This latter time interval corresponds better to the usual occurrence time of a second primary cancer in TP53 families [18].

In summary, this case report illustrates the difficulties in making a definitive diagnosis of LFS in patients lacking a positive family history and also the dangers of treating their primary tumors with radiotherapy when no proper genetic diagnosis has been made. In a near future, availability of high throughput technologies will allow more rapid genetic testing. Thus, patients at risk could be diagnosed earlier as carrier of detrimental gene mutations.

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