Plenary Paper

Dramatic efficacy of vemurafenib in both multisystemic and refractory Erdheim-Chester disease and Langerhans cell histiocytosis harboring the BRAFV600E mutation

*Julien Haroche,1,2 *Fleur Cohen-Aubart,1,2 *Jean-François Emile,3 *Laurent Arnaud,1,2 Philippe Maksud,4 Frédéric Charlotte,5 Philippe Cluzel,6 Aurélié Drier,7 Baptiste Hervier,1,2 Neïla Benamer,8 Sophie Besnard,9 Jean Donadieu,1,10 and Zahir Amoura1,2

1Department of Internal Medicine and French reference Center for Rare Auto-immune and Systemic Diseases, Assistance Publique–Hôpitaux de Paris (AP-HP), Pitié-Salpêtrière Hospital, Paris, France; 2Université Pierre et Marie Curie, UPMC University Paris 6, Paris, France; 3Research Unit (EA) EA4340 and Pathology Laboratory, Versailles University and AP-HP, Boulogne, France; Departments of 4Nuclear Medicine, 5Pathology, 6Radiology, 7Neuroradiology, and 8Pharmacy, Hôpital Pitié-Salpêtrière, University Paris 6, AP-HP, Paris, France, 9Department of Internal Medicine, Pontchaillou University Hospital, Rennes Cedex, France, and 10Department of Pediatrie, AP-HP, Centre de Référence des Histiocytes, Hôpital Trousseau, Paris, France

Key Points

- Treatment with vemurafenib induced a dramatic response in 3 patients with histiocytosis harboring BRAFV600E mutations.
- Tumor response was observed in both Erdheim-Chester disease and Langerhans cell histiocytosis.

Histiocytoses are rare disorders of unknown origin with highly heterogeneous prognosis. BRAFV600E gain-of-function mutations have been observed in 57% of cases of Langerhans cell histiocytosis (LCH) and 54% of cases of Erdheim-Chester disease (ECD), but not in other types of histiocytoses. Targeted therapy with an inhibitor of mutant BRAF (vemurafenib) improves survival of patients with melanoma. Here, we report vemurafenib treatment of 3 patients with multisystemic and refractory ECD carrying the BRAFV600E mutation; 2 also had skin or lymph node LCH involvement. The patients were assessed clinically, biologically (CRP values), histologically (skin biopsy), and morphologically (positron emission tomography [PET], computed tomography and magnetic resonance imaging). For all patients, vemurafenib treatment led to substantial and rapid clinical and biologic improvement, and the tumor response was confirmed by PET, computed tomography, and/or magnetic resonance imaging 1 month after treatment initiation. For the first patient treated, the PET response increased between months 1 and 4 of treatment. The treatment remained effective after 4 months of follow-up although persistent disease activity was still observed. Treatment with vemurafenib, a newly approved BRAF inhibitor, should be considered for patients with severe and refractory BRAFV600E histiocytoses, particularly when the disease is life-threatening. (Blood. 2013;121(9):1495-1500)

Introduction

Erdheim-Chester disease (ECD) is a rare non-Langerhans cell histiocytosis, characterized by the infiltration of tissues by foamy CD68+/CD1a− histiocytes.1 It is a systemic disease with diverse manifestations: the clinical course mainly depends on the extent and distribution of the disease, and ranges from asymptomatic bone lesions to life-threatening manifestations.2 Rare cases of ECD associated with Langerhans cell histiocytosis (LCH) have been reported.3

Unlike ECD, LCH histiocytes are CD1a+ and frequently infiltrate the epidermis in skin lesions.4 Interferon α (IFN) is generally the first choice for ECD therapy and improves survival. It should be prescribed at high dose if there is central nervous system and/or cardiovascular involvement.5 However, long-term IFN treatment can lead to severe side effects and some patients are refractory to treatment. Moreover some patients with CNS and/or cardiovascular infiltrations develop secondary resistance to high-dose of IFN. Alternative treatments include recombinant human interleukin-1 receptor antagonist, cladribine, thymidine kinase inhibitors, autologous hematopoietic stem cell transplantation.6 However, the optimal second line therapeutic strategy remains to be defined, mostly because these treatments have been evaluated in only small numbers of patients. Despite recent therapeutic progress the overall mortality remains high (18% of the 84 ECD patients seen at our institution). BRAFV600E mutations have been observed in 38% to 69% of cases of LCH.5-7 We recently reported BRAFV600E mutations in 54% of 24 patients with ECD.8 Vemurafenib, an inhibitor of mutant BRAF, has shown some efficacy against 2 diseases (melanoma and hairy-cell leukemia) associated with the BRAFV600E mutation.9,10

Methods

This study was approved by the ethics committee Ile de France III (#2011-A00447-34) and conducted in accordance with the Declaration of Helsinki.

Patient no. 1

A 65-year-old man presented in October 2010 with elevated serum creatinine and C-reactive protein (CRP) levels found on routine blood testing. Abdominal computed tomography (CT) disclosed retroperitoneal...
fibrosis complicated by bilateral hydronephrosis with sheathing of the abdominal aorta. Double ureteral pigtail stents were inserted and because hydronephrosis relapsed, bilateral nephrostomy was performed. Long-bone X-rays showed typical bilateral and symmetric cortical osteosclerosis in the lower limbs. A biopsy sample of the perirenal fibrosis showed numerous foamy CD68\(^+\) CD1a\(^-\) histiocytes. A diagnosis of ECD was established. Cerebral magnetic resonance imaging (MRI) showed sinus osteosclerosis; on cardiac MRI, the pericardium appeared thickened and there was specific infiltration of the auriculo-ventricular sulcus. Positron emission tomography (PET) revealed an intense uptake in the lower limbs and in the thoracic aorta.

Pegylated interferon α (PEG-IFN) treatment was started in July 2011, at 135 μg/w for 4 weeks, and was then increased to 180 μg/w due to cardiac involvement. Double ureteral pigtail stents were implanted to replace the nephrostomy and the blood creatinine concentration remained stable at 115 μM. In January 2012, the CRP level was 21 mg/L, and the creatinine level was 137 μM. The patient developed pruritus, and gammaglutamyl transferase (GGT) activity was 936 U/L (normal 12-55) and alkaline phosphatase activity was 644 U/L (normal 40-120). Abdominal CT showed the persistence of sheathing of the whole aorta. Bilateral hydronephrosis had increased and there was a left renal artery stenosis. Liver MRI and biopsy showed no or only minor abnormalities, and in particular no histiocytes or signs of sclerosing cholangitis. PET revealed that the number of lesions and uptake by the lesion had increased, that is, hypermetabolism of the long bones of the lower limbs, and of thoracic and abdominal aorta, mediastinum and retroperitoneum (Figure 1). Cardiac MRI findings were unchanged. Angioplasty of the left renal artery with stent implantation was performed. In February 2012, the IFN dose was increased (to 180 μg/w PEG-IFN alternating with 270 μg/w) due to the absence of response to treatment; this was followed by depression and fatigue. The depressive symptoms persisted despite administration of antidepressant drugs, leading to the IFN treatment being stopped in March 2012. The creatinine level increased to 190 μM, and consequently the double ureteral pigtail stents were replaced by thermoformable spiral metallic stents.

Patient no. 2

A 59-year-old woman was referred to us in February 2011 with a 2-year history of bone pain in the lower limbs. For the previous 6 months she had had diabetes insipidus, right orbital pain requiring morphine, and headaches. She had a history of bilateral breast cancer in 1998 treated by right mastectomy, radiotherapy, and chemotherapy. Thoracic and abdominal CT showed peri-aortitis (involving the arch, origin of the supra-aortic vessels, and the sub-renal aorta), nodular infiltration of the epiploon, and soft-tissue thickening of the pre- and retro-portal spaces. Cardiac MRI showed a right pseudo-tumoral atrial infiltration. Orbital MRI found a right retro-orbital infiltration and compression of the optic nerve with intense enhancement after gadolinium administration. Bone scintigraphy was highly suggestive of ECD with intense uptake in the lower limbs. A monoclonal IgA k component (12 g/L) was found, so a bone marrow biopsy was collected and revealed 30% plasma cell infiltration; renal function and calcemia were normal, hemoglobin was 12.5 g/dL, there was no proteinuria, and no lytic bone lesions were found. Stage I myeloma in the Durie-Salmon classification was diagnosed. Celioscopy in January 2011 found a peritoneal infiltration with foamy CD68\(^+\) CD1a\(^-\) histiocytes consistent with a diagnosis of ECD. At initial presentation, the patient had papulo-macular squamous intertrigo-like lesions under the left breast: biopsies showed Langerhans cell infiltrate in the epidermis and non-Langerhans cell infiltrate in the hypodermis (Figure 2). Initial PET revealed intense uptake in the retro-orbital space, right atrium, and abdominal aorta, and bilateral and symmetric uptake in the diaphyseal and metaphyseal regions of the long bones (Figure 1) typical of ECD. These findings led to a diagnosis of ECD associated with histologic lesions of LCH in the skin. The most severe symptoms, requiring therapy, were due to ECD (peri-aortitis, bone pain, pseudo-tumoral atrial infiltration, mesenteric localization, and right retro-bowel involvement). The associated IgAκ stage I myeloma only required monitoring.

Therapy with 135 μg/w PEG-IFN was initiated in March 2011. Partial improvement was observed, but pain in the right eye and cardiac abnormalities persisted. The PEG-IFN dose was increased to 180 μg/w in June 2011. In December 2011, the orbital pain had worsened and orbit MRI detected a...
new hypersignal in the right optic nerve; PET revealed intense uptake in the long bones. PEG-IFN, which had been maintained at 180 μg/w, was stopped at the beginning of March 2012 due to depression, fatigue, neutropenia (≈ 500/mm³), thrombocytopenia (46.000/mm³), and inefficacy against the retro-orbital involvement. The patient was re-assessed on March 23: she had fever, CRP was 89 mg/L, and PET confirmed the worsening of the retro-orbital, bone, and mediastinal involvements. Orbit MRI also showed persistence of the retro-orbital lesion (Figure 3). Treatment with 100 mg/d anakinra was initiated on March 24 and effectively reduced the fever but not the CRP value (Figure 4). Concomitantly, the cutaneous lesions relapsed leading to anakinra discontinuation on April 17. The right retro-orbital pain worsened. Skin lesions were re-biopsied and the findings were unchanged (co-existence of LCH in the epidermis and ECD in the hypodermis).

Patient no. 3
A 31-year-old woman originating from Laos was referred to our department in April 2010 for subacute renal failure, diabetes insipidus, and elevated CRP level. Physical examination identified bilateral superior and inferior eyelid xanthelasma. Blood tests confirmed elevated creatinine and CRP levels (178 μM and 147 mg/L, respectively). Abdominal CT-scan revealed retroperitoneal fibrosis and lymphadenopathy, and cerebral MRI showed that the pituitary axis was enlarged. Long bone 99Tc scintigraphy showed intense metaphyseodiaphyseal uptake in femurs and tibias. Similar hypermetabolism in the long bones and also in the spleen, and retroperitoneal adenopathies were found on PET.

Biopsy of the palpebral xanthelasma confirmed the ECD localization with foamy histiocytes. Immunohistology was positive for CD68 and negative for PS100 with a few CD1a-positive cells. Biopsies of T4 vertebra and right inferior metaphyseal femur showed the same ECD patterns, and a biopsy of a retroperitoneal node displayed characteristics of LCH with strongly positive CD1a and PS100 staining.

In April 2010, treatment with PEG-IFN (135 μg/w) was started and double ureteral pigtail stents were inserted. Ten months later, in February 2011, the CRP level was 24 mg/L, creatinine had decreased to 138 μM, and PET showed an improvement of long-bone uptake. In February 2012, the clinical status had deteriorated with new lower-limb pain and increased size of xanthelasmas. Abdominal CT scan showed enlargement of periaortic and periureteral infiltrations. Cardiac MRI, which had been normal on previous assessments, displayed a typical infiltration of the right atrial wall, atrioventricular groove, and gadolinium enhancement of the left ventricular wall, septum, and apex. The CRP level was 20 mg/L and PET disclosed a strong, new uptake in the basilar skull region. The PEG-IFN dose was increased to 180 μg/w on March 20, but was clinically ineffective: the CRP level increased to 42 mg/L. Treatment was stopped on May 9.

Histology and molecular analyses
All biopsies (perirenal, peritoneal, skin, and bone) obtained from the 3 patients before or after vemurafenib treatment were reviewed by 2 pathologists with specialist training in histiocytosis (F.C.-A., J.-F.E.). All patients provided consent for vemurafenib treatment after the provision of counseling. Each sample was processed for immunohistochemistry with antibodies against CD1a (Beckman-Coulter), CD68 (Dako), CD163 (Thermoscientific) and S100 protein (Dako), and with VE1 (a mouse monoclonal antibody specific for the BRAFV600E mutant, kindly provided by Prof A. von Deimling, Heidelberg University, Heidelberg, Germany). Clone L26 (anti-CD20) was used as IgG2a isotype matched control for the VE1 antibody.11 BRAFV600E mutations were detected as previously described.12 DNA was extracted from formalin-fixed and paraffin-embedded (FFPE) tissues after histologic detection of histiocyte-rich areas, and pyrosequencing with PyroMark Q24 (QIAGEN) was used to detect BRAFV600E mutations.

Results
BRAFV600E mutations were detected in ECD biopsies from all 3 patients, and in samples of LCH epidermal infiltration from...
The expression of the V600E mutant BRAF protein in histiocytes from both ECD and LCH lesions was confirmed by immunohistochemistry (Figure 2).

The efficacy of vemurafenib

These 3 cases of refractory ECD with life-threatening manifestations associated with the \( \text{BRAF}^{\text{V600E}} \) mutation were all treated with vemurafenib. All patients provided consent for vemurafenib treatment after appropriate counseling.

Vemurafenib (initially given at 1920 mg/d) was tapered in all 3 patients to 960 mg/d from day 30 due to erythema, and was thereafter maintained at this dose. Within a few days of the initiation of treatment, itching disappeared. Evaluation on day 30 showed an improvement of creatinine (190 to 161 \( \mu \)M), GGT (936 to 300 U/l) and alkaline phosphatase (644 to 177 U/l) values. On day 62, GGT was 183 U/l, PAL 146 U/l, and creatinine 170 \( \mu \)M. CRP was normalized by day 30 (Figure 4), and remained under 5 mg/l on day 62. PET was performed after 30 days of vemurafenib treatment and the findings compared with those obtained 4 months earlier: there was a substantial improvement of all lesions (Figure 1), with a mean Standardized Uptake Value (SUV) change of \(-70\%\) (range, \(-66\%\) to \(-80\%\)); the change within soft tissue and bone was similarly \(-70\%\) (range, \(-51\%\) to \(-78\%\); supplemental data, available on the Blood Web site; see the Supplemental Materials link at the top of the online article). Thoraco-abdominal CT after 37 days of treatment showed a significant reduction of periaortic infiltration (Figure 3). This cannot be attributed to PEG-IFN treatment, withdrawn on March 31, 2012, because several patient no. 2. showed clear clinical improvement within a few days of initiation of this treatment.

**Patient no. 1.** Vemurafenib was started at 1920 mg/d (b.i.d.) on April 19, 2012 and was tapered to 960 mg/d from day 30 due to erythema, and was thereafter maintained at this dose. Within a few days of the initiation of treatment, itching disappeared. Evaluation on day 30 showed an improvement of creatinine (190 to 161 \( \mu \)M), GGT (936 to 300 U/l) and alkaline phosphatase (644 to 177 U/l) values. On day 62, GGT was 183 U/l, PAL 146 U/l, and creatinine 170 \( \mu \)M. CRP was normalized by day 30 (Figure 4), and remained under 5 mg/l on day 62. PET was performed after 30 days of vemurafenib treatment and the findings compared with those obtained 4 months earlier: there was a substantial improvement of all lesions (Figure 1), with a mean Standardized Uptake Value (SUV) change of \(-70\%\) (range, \(-66\%\) to \(-80\%\)); the change within soft tissue and bone was similarly \(-70\%\) (range, \(-51\%\) to \(-78\%\); supplemental data, available on the Blood Web site; see the Supplemental Materials link at the top of the online article). Thoraco-abdominal CT after 37 days of treatment showed a significant reduction of periaortic infiltration (Figure 3). This cannot be attributed to PEG-IFN treatment, withdrawn on March 31, 2012, because several
other markers clearly indicated that the patient was a nonresponder (high CRP values, hydrenephrosis and pyloric infiltration, worsening of PET findings). Hydrenephrosis and pyloric infiltration resolved completely. Note that the PET assessment at 4 months showed that the decrease of FDG uptake had been sustained: uptake was lower than that observed after the first month (Figure 1). Aortic MRI after 4 months disclosed a marked regression of peri-aortic sheathing relative to that observed by CT-scan in May 2012. The CRP value remained low on day 123 of treatment (Figure 4).

**Patient no. 2.** Vemurafenib was administered at 1920 mg/d (b.i.d) from April 17 with a transient interruption of 2 days (days 12 to 14) due to headaches and fever at 39°C. The CRP value increased to 114 mg/L (May 1); lumbar puncture was normal, blood cultures were negative, and both physical examination and thoracic CT-scan ruled out pneumopathy. The clinical symptoms rapidly disappeared and CRP values declined (100 and 48 mg/L on May 2 and 3). We were able to resume treatment at full dose at day 14. Our impression was that this transient episode was perhaps a viral infection, or cytokine-related, but not due to the disease or vemurafenib treatment. The dose was tapered to 960 mg/d from day 30 due to itching, skin rash and keratosis pilaris lesions which had appeared a few days before. Itching was not attributed to LCH involvement but appeared to be a consequence of the cutaneous toxicity of vemurafenib. Skin manifestations disappeared within days of the reduction of the dose which was maintained at 960 mg/d until further follow-up.

After a few days, histiocytic skin lesions resolved (Figure 2), and the pain in the right eye disappeared such that morphine could be stopped. A new skin biopsy confirmed the absence of histiocytic infiltration of the epidermis and hypodermis. CRP was normalized by day 30 (Figure 4), and was 5.2 mg/L (N < 5) on day 48. PET 31 days after initiation of vemurafenib was compared with that performed 2 months earlier: the change in the mean SUV was −57% (range, −51% to −67%) and that in soft tissue and bone was −70% (range, −56% to −78%; Figure 1 and supplemental data). Orbit MRI on day 36 showed a 26% to 66% decrease of the retro-orbital infiltration (Figure 3 and supplemental data). Thoraco-abdominal CT on day 39 demonstrated a 30% to 40% decrease of the posterior and lateral peri-aortic infiltration (Figure 3), and the absence of the previously observed interlobular septa thickening, characteristic of pulmonary involvement of ECD. The CRP value remained below 5 mg/L on day 141 of treatment (Figure 4), and the patient was asymptomatic, with the absence of orbital pain.

**Patient no. 3.** Vemurafenib was administered at 1920 mg/d (b.i.d) from May 9 to May 29: the dose was tapered to 960 mg/d (b.i.d) due to pilar keratosis and erythema. One month later, the CRP level was 10 mg/L, the thickness of xanthelasmas had diminished, and PET displayed significantly decreased uptake in the basilar region and by the long bones. In July 2012, the CRP value was normal. The most striking feature was the significant improvement of the cardiac MRI on July 19: right atrial wall infiltration had regressed substantially (Figure 3). On day 119 of treatment, the CRP value was still low, at 8 mg/L (Figure 4).

### Discussion

This is the first report of the use of vemurafenib in ECD patients carrying the BRAFV600E mutation, with a multisystemic form of the disease refractory to IFN. The treatment was extremely and rapidly effective in all 3 patients: clinical symptoms improved, CRP values normalized, all pathologic uptakes on PET regressed substantially, peri-vascular sheathing regressed, and heart infiltration improved markedly; the LCH skin lesions of patient no. 2 disappeared and the severity of the orbital lesion decreased.

PET assessment of treatment of cases of ECD has not previously documented such rapid efficacy. IFN treatment has a much more subtle and slow effect on the regression of uptakes. For patient no. 1, there was a mean regression of 70% of visceral and bone 18F-fluorodeoxyglucose uptakes at month 1, and the therapeutic response observed at month 4 was even better. Substantial regression of peri-vascular sheathing and disappearance of hydrenephrosis (CT), as well as normalization of measures of CRP and liver enzymes confirmed the excellent therapeutic efficacy. The benefits of treatment were similar in patient no. 2, with 57% to 70% regression of visceral and bone 18F-fluorodeoxyglucose uptakes, regression of peri-aortic and orbital infiltrations, and normalization of CRP values.

Imaging and analysis of biologic markers demonstrated tumor regression within 40 days in patients 1 and 2, and for patient no. 3 there was a major regression of heart infiltration by day 71. Moreover, the clinical benefits, with the resolution of pruritus for patient no. 1 and of skin lesions and orbital pain for patient no. 2, were apparent within a few days. The Pitie-Salpetriere Hospital has cared for 84 ECD patients between 1991 and 2012, and there have been no previous cases of such remarkable and rapid clinical, biologic, and imaging improvements.

Typical LCH infiltration has been reported in some patients with confirmed ECD. In patient no. 2, the BRAFV600E mutation was detected in both types of histiocytes. LCH skin lesions disappeared rapidly under vemurafenib and this was confirmed by biopsy, suggesting that LCH patients may also benefit from vemurafenib treatment; the value of such treatment should be assessed in patients with pure LCH disease.

Contrasting with other types of histiocytosis, BRAFV600E mutations are present in approximately half of the patients with ECD or LCH. Among the 84 ECD patients followed in our center, 17 (20.2%) have died of disease. Although an appropriate dosing schedule and treatment duration remain to be determined in clinical trials, our findings provide compelling evidence of the fast efficacy of BRAF inhibition in BRAFV600E-associated ECD. The long-term efficacy of BRAF inhibition in ECD should also be studied, as secondary resistance develops in almost all cases of BRAFV600E-associated melanoma. Patient no. 1 was asymptomatic at 4 months and had normal CRP values. At that time, follow-up PET assessment only revealed persistent bone uptake, which we believe should rather be interpreted as being a typical hallmark of the disease than a marker of disease activity. Nevertheless, the durability of the therapeutic response to vemurafenib remains unknown and should be evaluated over longer periods.

In view of the remarkable efficacy in the 3 cases we report, we believe that BRAF inhibition should be considered in a larger cohort of BRAFV600E-associated histiocytosis patients, particularly those with life-threatening disease.

### Acknowledgments

The authors thank Prof A. von Deimling (Heidelberg, Germany) for providing the anti-BRAFV600E mouse monoclonal antibody VE1.

This work was supported in part by grants from the nonprofit organism Association pour la Recherche en Pathologie (AREP).
**Authorship**


**Conflict-of-interest disclosure:** J.H. received honoraria from Glaxo Smith Kline for counseling of patients with histiocytosis on the treatments with targeted therapies. J.-F.E. received honoraria from Roche and Glaxo Smith Kline for counseling patients with melanomas on the diagnosis and/or treatment with BRAF inhibitors. The remaining authors declare no competing financial interests.

**Correspondence:** Dr Julien Haroche, MD, PhD, Service de Médecine Interne 2, Groupe Hospitalier Pitié-Salpêtrière, 47-83 bd de l’Hôpital, 75013, Paris, France; e-mail: julien.haroche@psl.aphp.fr.

**References**

Dramatic efficacy of vemurafenib in both multisystemic and refractory Erdheim-Chester disease and Langerhans cell histiocytosis harboring the BRAF V600E mutation

Julien Haroche, Fleur Cohen-Aubart, Jean-François Emile, Laurent Arnaud, Philippe Maksud, Frédéric Charlotte, Philippe Cluzel, Aurélie Drier, Baptiste Hervier, Neïla Benameur, Sophie Besnard, Jean Donadieu and Zahir Amoura