Conclusions Current calciphylaxis treatments alternatives aim to lower the serum calcium phosphate concentration thereby preventing, or even reversing, calcium phosphate oversaturation, precipitation and, finally, calcification. Administration of IV sodium thiosulfate, which sequesters calcium ions to form highly soluble calcium thiosulfate complexes, can prevent calcium phosphate precipitation.

No conflict of interest.


doi:10.1136/ehjphp-2013-000276.603

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Background Glioblastoma multiforme (GBM) are primary brain tumours that are currently incurable. Despite a well codified first-line treatment with concomitant radio-chemotherapy (temozolomide), recurrences of GBM occur and have limited treatment options. Furthermore, there is a lack of effective therapies and no standard relapse treatment. Anti-angiogenic drugs, such as bevacizumab, show encouraging results for patients with recurrences of high-grade gliomas.

Purpose To describe treatments of GBM relapses within three cities in northeast France. Nancy, Reims and Strasbourg. We especially tried to assess the impact of bevacizumab on survival endpoints.

Materials and Methods This is a retrospective study with GBM patients diagnosed between 2006 and 2008. Medical data describing the population and therapeutic oncology support were collected in each site from individual patient charts. Overall Survival (OS) and Progression Free Survival (PFS) were estimated by the Kaplan-Meier method and compared by the log-rank test.

Results Between 2006 and 2008, 521 patients were diagnosed with GBM, of whom 135 patients were treated for at least one recurrence. There were 95 males and 38 females; median age at diagnosis was 58 years. Main relevant signs of the initial tumour were intracranial hypertension and epilepsy. Initial treatment consisted for 64% of patients in surgical excision, and 86% of patients received conventional radio-chemotherapy followed by adjuvant temozolomide.

More than 50% of recurrences were diagnosed on both clinical and radiological grounds. Discarding palliative care, almost all patients with GBM relapse received chemotherapy: 95% at first recurrence (n = 126/135), 95% at second recurrence (n = 67/73) and 100% at third recurrence (n = 26/26). Bevacizumab was used (alone or in association) in a third to half of cases.

In our population, neglecting the type of relapse treatment, median OS was 17.8 months [5–50 months]. When patients received bevacizumab at some point in their care, median OS was 20.2 months [7–90 months]. This OS is significantly different from the median OS observed without bevacizumab which was 13.5 months [5–41 months]. PFS until the second recurrence with bevacizumab was 5.5 months compared to 3.1 months without bevacizumab. PFS until the third recurrence with bevacizumab was 5 months against 2 months without bevacizumab. However, these results do not show bevacizumab providing significant PFS improvement, especially in the long term.

Conclusions Bevacizumab seems to improve OS in patients with GBM recurrences. However only prospective randomised studies will define the appropriate strategy treatment in recurrent glioblastoma. This work is one of the several projects of the NENO group which aims to standardise practise and build rational standards.

No conflict of interest.

TREATMENT OF HEPATIC METASTASES FROM MELANOMA WITH IRINOTECAN LOADED IN ELUTING BEADS

doi:10.1136/ehjphp-2013-000276.604

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Background Chemoembolization of hepatic melanoma metastases refractory to treatment using irinotecan-loaded DC beads [embolic Drug-Eluting Beads]: a novel palliative treatment with which there is as yet little experience.

Purpose To show the progress of a clinical case of metastatic choroidal melanoma treated with irinotecan-loaded DC Beads.

Materials and Methods The pharmacy department loaded the particles with irinotecan ourselves and monitored the patient through the clinical history. The patient was a 38-year-old man with stage IV choroidal melanoma in the left eye (2007).

Results In October 2011, 4 hepatic nodules were detected: 3 in segment VII (23, 25, 11 mm) and 1 in segment II (16 mm), 2 cycles of dacarbazine treatment (1649 mg × 1day) stabilised the disease. The patient experienced emesis and diarrhoea. Given this intolerance and negative BRAFV600E mutation, ipilimumab reinforcement treatment was administered (225 mg × 1day q21days). After 4 cycles of ipilimumab, the disease stabilised for 5 months. In May 2012, an increase in size of the nodules was described and 6 new nodules in both hepatic lobes: segment II (42 × 34 mm), IVb (15 mm), VII (25, 26 and 61.4 × 48 mm) and VIII (14 mm) were observed. Surgery was rejected due to the presence of multinodular lesions and transarterial chemoembolization with irinotecan-loaded DC beads was attempted.

Hypervascular lesions were observed in the distal branches of the hepatic artery by bilar hepatic arteriography using selective catheterization of both hepatic arteries. Subsequently, hepatic chemembolization was performed by administering 100 mg irinotecan-loaded beads (75–100 microns). After 2 cycles in each hepatic lobe, treatment response was assessed by the RECIST criteria. One month after the last chemoembolization, stable disease (no new nodules and arterial necrosis <30%) was confirmed. No immediate complications were observed, except for a slight elevation of hepatic enzymes that resolved.

Conclusions Hepatic chemoembolization using irinotecan-loaded beads is a viable alternative with good prognosis for hepatic metastases of choroidal melanoma. A higher concentration of chemotherapeutic drug is achieved within the hepatic lesions using lower concentration of chemotherapy in others hepatic lesions, improving clinical results.

No conflict of interest.

TREATMENT OF SEVERE PSORIASIS WITH BIOLOGICALS

doi:10.1136/ehjphp-2013-000276.605

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Background Biological drugs are a relatively new class of treatment for severe psoriasis (SP).

Purpose To analyse the use and outcomes of biologicals in SP.

Materials and Methods Retrospective observational study for 23 months of patients with SP who had not previously received biologic treatment.

No conflict of interest.
biological therapy (treatment-naive). Data were collected from the pharmacy outpatient dispensing programme and the clinical history.

Results 46 treatment-naive patients started treatment, mean age 43 (17–83), 58.7% men. They were treated with: adalimumab (20 patients), infliximab (17) and ustekinumab (9); none with etanercept.

Abstract CPC-148 Table 1

<table>
<thead>
<tr>
<th></th>
<th>Adalimumab</th>
<th>Infliximab</th>
<th>Ustekinumab</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lost to follow-up</td>
<td>5% (1/20)</td>
<td>23.5% (4/17)</td>
<td>22% (2/9)</td>
</tr>
<tr>
<td>Continuing treatment</td>
<td>40% (8/20)</td>
<td>29.4% (5/17)</td>
<td>67% (6/9)</td>
</tr>
<tr>
<td>Withdrawal/</td>
<td>5% (1/20)</td>
<td>5.9% (1/17)</td>
<td>11% (1/9)</td>
</tr>
<tr>
<td>Discontinuation by doctor</td>
<td>20% (4/20)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Change of treatment</td>
<td>30% (6/20)</td>
<td>41.2% (7/17)</td>
<td>–</td>
</tr>
</tbody>
</table>

The causes of the discontinuation/change of treatment were:

Abstract CPC-148 Table 2

<table>
<thead>
<tr>
<th></th>
<th>Adalimumab</th>
<th>Infliximab</th>
<th>Ustekinumab</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lack of efficacy</td>
<td>27.2% (3/11)</td>
<td>25% (2/8)</td>
<td>–</td>
</tr>
<tr>
<td>Adverse reactions</td>
<td>18.2% (2/11)</td>
<td>37.5% (3/8)</td>
<td>–</td>
</tr>
<tr>
<td>Lack of response + adverse reactions</td>
<td>18.2% (2/11)</td>
<td>12.5% (1/8)</td>
<td>–</td>
</tr>
<tr>
<td>Lack of adherence</td>
<td>–</td>
<td>25% (2/8)</td>
<td>–</td>
</tr>
<tr>
<td>Good response</td>
<td>36.4% (4/11)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Other reasons</td>
<td>–</td>
<td>–</td>
<td>100% (1/1)</td>
</tr>
</tbody>
</table>

Adverse reactions that caused withdrawal or change were:

Abstract CPC-148 Table 3

<table>
<thead>
<tr>
<th></th>
<th>Adalimumab</th>
<th>Infliximab</th>
<th>Ustekinumab</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autimmune hepatitis</td>
<td>25% (1/4)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Asthenia and mood changes</td>
<td>25% (1/4)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Psoriatic arthritis</td>
<td>25% (1/4)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Pharyngitis and candidiasis</td>
<td>25% (1/4)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Acute infusion reactions</td>
<td>75% (3/4)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Psoriatic arthropathy</td>
<td>25% (1/4)</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

Conclusions The first biological in treatment-naive patients was 1st) adalimumab, 2nd) infliximab and 3rd) ustekinumab. Ustekinumab was the biological drug that achieved the best retention rate. Several patients discontinued their treatment with adalimumab because of good response, since it can be used in intermittent treatment schemes. Change in treatment was more frequent with infliximab, mainly because of infusion reactions. Ustekinumab was the only biological that didn’t cause adverse reactions that caused withdrawal or change.

No conflict of interest.

CPC-149 TRIMEBUTINE: A CASE OF ABUSE AND POSSIBLE DEPENDENCE
doi:10.1136/ehjpharm-2013-000276.606

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Background Trimebutine has an agonist effect on digestive tract mu, kappa, and delta opiate receptors.

Purpose To describe a case report of an abuse and a possible dependence on trimebutine.

Materials and Methods Medical record review and literature search about trimebutine dependence.

Results A 46-year-old woman with a history of Chronic Intestinal Pseudo-Obstruction (CIPO) was prescribed amikacin and trimebutine in the hospital since 2011. Her gastroenterologist initially prescribed trimebutine at 100 mg intravenously three times a day, with a possibility of 100 mg shots if necessary without a maximum dose. At the same time she obtained another prescription by her general practitioner (50 mg IV if needed). Finally 735 ampoules were delivered in seven weeks (15 a day). This overconsumption alarmed the pharmaceutical team and a literature review was made. Dependence is described in a French register: six cases of intravenous abuse or dependence were reported between 1993 and 2009. At high doses trimebutine is cardiotoxic (bradycardia, rhythm disorders) and neurotoxic (convulsions). We alerted the prescribers and reported this abuse to our pharmacovigilance centre. A questioned to evaluate the level of dependence was sent to the general practitioner.

Once the general practitioner had been informed, the gastroenterologist alone managed her CIFO treatment and a new prescription was established with a trimebutine posology more consistent with the marketing authorization.

Conclusions Provision from a hospital enabled us to detect the overuse of this drug. Dependence is difficult to prove and drug abuse screening test in the assessment of DSM IV should be used to establish it.

No conflict of interest.

CPC-150 TUBERCULOSIS AND SYSTEMIC DISEASES
doi:10.1136/ehjpharm-2013-000276.607

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Background The immunosuppression of systemic diseases makes the management of patients with tuberculosis more complicated.

Purpose To monitor the clinical evolution of tuberculosis in patients suffering from systemic diseases.

Materials and Methods A retrospective study, from 1998 to 2012, in the internal medicine service in Rabta hospital, Tunisia, of 9 patients (8 women and a man, median age: 54 years) suffering from connective tissues disease, treated by corticosteroids linked in one or several treatments to immunosuppressants, who subsequently developed tuberculosis.

Results The median time to diagnosis was 116 days (7d – 1 year). The location of the tuberculosis was pulmonary (n = 2), ganglionic (n = 3), urogenital (n = 2), tubercular spondylodiscitis (n = 1), more than one location (n = 1). The diagnosis of tuberculosis was confirmed by bacteriology (n = 4) four cases, histologically (n = 5) and by a test treatment (n = 1). Systemic illnesses were: systemic lupus erythematous (n = 5), Gougerot-Sjögren syndrome (secondary or primary) (n = 3), sarcoidosis (n = 1), systemic sclerosis linked to pernicious anaemia (n = 1), rheumatoid arthritis (patient 2 linked to lupus) (n = 1) and multiple auto-immune syndrome (n = 1). The diagnosis of systematic illness was made before that of tuberculosis in 8 patients and concomitantly in only one. Under treatment by four drugs then by two drugs, the evolution of tuberculosis was favourable in five of nine patients. One of the patients developed an allergy in isoniazid and resistance to the anti-tubercul treatment. Five of our patients recovered from their illness.

Conclusions This study confirms the often extra-pulmonary character of tuberculosis in patients with systemic disease as well as the difficulty of diagnosis and problems multiplied by this association.

No conflict of interest.

CPC-151 TYPE OF CANCER AND RISK FACTORS IN HIV PATIENTS ON ANTRETROVIRAL TREATMENT
doi:10.1136/ehjpharm-2013-000276.608

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CPC-148 Treatment of Severe Psoriasis with Biologicals

J Ruiz Gutiérrez, G Roustán Gullón, P Calabuig Martínez and A Torralba Arranz

_Eur J Hosp Pharm_ 2013 20: A218-A219
doi: 10.1136/ehjpharm-2013-000276.605

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