Cerebral venous thrombosis: a Moroccan retrospective study of 30 cases

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Abstract

Cerebral venous thrombosis (CVT) is a rare origin of stroke, the clinical presentation and etiologies vary. The prognosis is shown to be better than arterial thrombosis. Magnetic Resonance Imaging (MRI) and MR Venography (MRV) are currently important tools for the diagnosis. We studied 30 cases of CVT diagnosed in the department of neurology at the University Hospital of Fez (Morocco). Patients diagnosed with CVT signs between January 2003 and October 2007 were included in the study. Cerebral CT-scan was performed in 27 cases (90%) while the MRI examination was done in 18 patients (67%); and most patients (90%) received anticoagulant therapy. The mean age of our patients was of 29 years (age range between 18 days and 65 years). A female predominance was observed (70%). The clinical presentation of patients was dominated by: headache in 24 cases (80%), motor and sensory disability in 15 cases (50%), seizures in 10 cases (33%), consciousness disorder in 10 cases (33%). CVT was associated to post-partum in 10 cases (33%), infectious origin in 8 cases (26%), Behçet disease in 2 cases (7%), pulmonary carcinoma in 1 case, thrombocytemia in 1 case and idioopathic in 7 cases (23%). The evolution was good in 20 cases (67%), minor squelaes were observed in 6 patients (20%), while major squelaes were observed in 2 cases. Two cases of death were registered. The CVT is a pathology of good prognosis once the diagnosis is promptly established and early heparin treatment initiated.
Introduction

CVT is a rare origin of stroke. This pathology was considered an infectious disease for a long period. In addition, it was assumed to be affecting the superior sagittal sinus dragging to death. Anticoagulants were contraindicated while the diagnosis was established at the autopsy stage. Currently CVT is a disease with various clinical presentations and etiologies. MRI and MRV are the key diagnosis tools. Heparin is the first-line treatment. The prognosis is better compared to arterial thrombosis with mortality well below 10%. The goal of this study is to describe and discuss different features of this pathology including frequency, clinical model, diagnosis and outcome.

Methods

We studied 30 cases colligated in the department of neurology of the University Hospital of Fez (Morocco). Patients were recruited from January 2003 to October 2007 (58 months). All patients demonstrating clear radiological signs of CVT were included. The cerebral CT-scan was performed in 27 patients (90%). The period between onset and CT scan varied from 1 day to 45 days. The MRI examination was achieved in 17 patients (56%). Ninety percent of patients (90%) received anticoagulant therapy. Considering the CT scan without injection of contrast agents, the inclusion criteria was spontaneous hyperdensity or cord sign; while contrast agents was injected the empty delta sign and empty lateral sinus sign were the inclusion criteria. On MRI, inclusion criteria were hypersignal on T1 and T2 sinuses (Figure 1), abnormal defect or rupture of the sinus in MRV except two cases of deep cerebral venous thrombosis who presented indirect signs of thrombosis of deep veins. The MRI T1 and T2 weighted were done in axial, sagittal and coronal slices. Axial FLAIR and echoplanar susceptibility-weighted (T2*) and T1 with gadolinium injection were also achieved. Our patients have had hyperleucocytosis in 8 cases (27%), raised erythrocyte sedimentation rate in 11 cases (37%), thrombophilia was recorded in four patients including low S-protein in three cases and C-protein in one case. The CT-scan showed direct signs such as empty lateral sinus in 11 cases (37%) (Figure 3), cord sign in six cases (20%), empty delta sign in seven cases (23%). The indirect signs were cortical hypodensity in 10 cases (33%), subcortical hypodensity in 12 cases (40%), parenchymal hematoma (1 case), enhancement of wall sinuses after contrast agent injection in 9 cases (30%), multiple and grouped hyperdensities in six cases (20%). 14 patients benefited both CT scan and MRI examinations. The CT scan performed diagnosis in five cases (35%), confirmed by MRI associated with MRV. The CT scan was normal in three cases (21%). The MRI associated with MRV was more efficient in demonstrating the lesion compared to CT in 9 patients (65%) (Table 2). The CT-scan and/or MRI including the MRV showed the CVT in the sagittal superior sinus (SSS) in 15 cases (50%), right lateral sinus (RLS) in 12 cases (40%), left lateral sinus (LLS) in 3 cases (10%), cavernous sinus (CS) involvement in 5 cases (16%), deep cerebral vein in two cases (7%), and cortical cerebral vein in one case. The diagnosis of cortical cerebral vein thrombosis was also demonstrated as a hypersignal on echo-planar susceptibility weighted (T2*) images (Figure 4). The diagnosis of deep cerebral vein thrombosis was performed by indirect signs on MRI, and theses cases have shown normal deep venous system on MRV images.

Etiology

All patients underwent systematically an investigation of the thromboembolic antecedent, clinical examination, blood cells enumeration, prothrombin time (PT), activated partial thromboplastin time (APTT), erythrocyte sedimentation rate (ESR) and a thoracic x-ray. The post-partum was the most frequent cause of CVT in our series consisting of 10 cases (33%); infectious origin was seen in 8 cases (26%), four cases of otomastoiditis, three cases of severe staphylococcal of the face and these patient have had delayed treatment, one case of ethmoiditis, one case of miliary tuberculosis; Behcet disease in 2 cases (7%); pulmonary carcinoma in 1 case; essential thrombocytopenia in 1 case and thombophilia in one case (deficiency in protein C) (Figure 5). Finally, in 7 cases (23%), the etiology of the CVT could not be established.

Results

The mean age of our patients was 29 years old, ranging from 18 days to 65 years old. High predominance of female involvement was noticed with sex ratio of 2.3. The average period between the onset of symptoms and consultation was 18 days. The Table 1 summarizes patients' data.

Clinical presentation

The mode of onset of symptoms was acute in 50% of cases, a subacute in 47% and progressive in one case (3%). The clinical presentation was variable, but the most common symptoms were headaches in 24 cases (80%), motor and sensitive disability in 15 cases (50%), seizures in 10 cases (33%) including two cases of status epilepticus, consciousness disorder was noticed in 10 cases (33%) and fever (40%) (Figure 2).

Biological and Radiological Examinations

The biological examination showed anemia in 13 cases (43%), hyperleucocytosis in 8 cases (27%), raised erythrocyte sedimentation rate in 11 cases (37%), thrombophilia was recorded in four patients including low S-protein in three cases and C-protein in one case. The CT-scan showed direct signs such as empty lateral sinus in 11 cases (37%) (Figure 3), cord sign in six cases (20%), empty delta sign in seven cases (23%). The indirect signs were cortical hypodensity in 10 cases (33%), subcortical hypodensity in 12 cases (40%), parenchymal hematoma (1 case), enhancement of wall sinuses after contrast agent injection in 9 cases (30%), multiple and grouped hyperdensities in six cases (20%). 14 patients benefited both CT scan and MRI examinations. The CT scan performed diagnosis in five cases (35%), confirmed by MRI associated with MRV. The CT scan was normal in three cases (21%). The MRI associated with MRV was more efficient in demonstrating the lesion compared to CT in 9 patients (65%) (Table 2). The CT-scan and/or MRI including the MRV showed the CVT in the sagittal superior sinus (SSS) in 15 cases (50%), right lateral sinus (RLS) in 12 cases (40%), left lateral sinus (LLS) in 3 cases (10%), cavernous sinus (CS) involvement in 5 cases (16%), deep cerebral vein in two cases (7%), and cortical cerebral vein in one case. The diagnosis of cortical cerebral vein thrombosis was also demonstrated as a hypersignal on echo-planar susceptibility weighted (T2*) images (Figure 4). The diagnosis of deep cerebral vein thrombosis was performed by indirect signs on MRI, and theses cases have shown normal deep venous system on MRV images.

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The treatment

The treatment of intracranial hypertension was based on carbonic anhydrase inhibitor or mannitol (53%), and the lumbar puncture was performed to remove the cerebrospinal fluid (23%). Most of our patients (90%) received heparinotherapy (low-molecular-weight heparin 68% or standard heparin 32%) followed by oral anticoagulant for 6 months. Four patients (10%) did not receive heparin, two have had a thrombosis of the cavernous sinus complicated by a carotid aneurysm; hence the anticoagulation was contraindicated. The third patient was a newborn of 18 days.

Antibiotherapy was administrated to patients with infectious origin (47%). Antiepileptics (27%) were prescribed for cases of seizures. We used steroids (prednisone 1mg/kg/day) in two cases of Behcet disease. Allopurinol (300mg/day) and hydroxycurvamin (1000mg/day) were used in cases of essential thrombocytemia. Two patients with severe disorder of consciousness were hospitalized in reanimation department; one of them has died in this department. The surgery was not the treatment of choice in any of the reported cases.
The outcome
After treatment, a full recovery was noticed in 18 cases (60%). Sequelae were observed in 8 patients (27%), these consisted of blindness in one case associated to cavernous sinus thrombosis complicated by severe staphylococcal of the face. Hemiparesis was noticed in 3 cases, tetraplegia associated to cognitive dysfunction was recorded in case of deep cerebral veins thrombosis. Epilepsy was noticed in a case stabilized using carbamazepin; ophthalmoplegia after a cavernous sinus thrombosis; pyramidal hypertonia in the newborn of 18 days. Three patients died (10%) including one patient in the post partum, one of cavernous sinus thrombosis and one idiopathic.

Discussion

Cerebral venous thrombosis is a rare type of stroke that might occur at any age. The diagnosis of CVT requires a visualization of the thrombosis, which is commonly demonstrated by cerebral MRI and MRV. The CT-scan might be helpful but does not allow establishing the diagnosis.

CVT affects about five people per million per year and represents 0.5 % of all stroke [1]. CVT might occur in any stage of life. Most authors agree on the predominance of CVT in young subjects, this was also revealed in our study. Similar agreement was found for the gender ratio [1]. In our series, we found a sex ration of 2.3. The female predominance is explained by the high frequency of CVT in the post-partum. The clinical presentation is determined by the patient age, the delay between the onset and the hospitalization, the location of the thrombosis and the occurrence of cerebral parenchymal lesions.

Headaches represent the main and first symptom of CVT retrieved in most studies. CVT with isolated headache is possible, without intracranial hypertension, subarachnoid haemorrhage, or meningitis signs and symptoms[2, 3]. Other symptoms are partial and generalized seizures, motor or sensitive deficits, altered consciousness, and papilloedema which might be associated to other symptoms [4]. Motor or sensitive deficit, altered consciousness and seizures are frequent in our study and agrees with earlier study (Einhäupl 1990), this was mostly related to delayed patients consultation. The 18 days newborn patient has shown a generalized seizures and hyporeactivity as main symptoms of the SSS thrombosis. The case number 20 has shown isolated cortical cerebral vein thrombosis revealed by motor deficit without seizures association.

In SSS thrombosis, signs are dominated by the unilateral or bilateral motor deficit seizures and consciousness disorder. On the other hand, the isolated intracranial hypertension syndrome is not frequent in SSS thrombosis. However, CVT of the LS is often revealed by an isolated intracranial hypertension syndrome. When the left LS is affected, an aphasia occurs, thereafter, a patient can present with Wernicke aphasia and partial seizures.

The clinical manifestations of deep CVT are more severe, it often presents with coma and bilateral motor deficit. These manifestations have been noticed in case number 18, while the case number 29 presented only a unilateral motor deficit. Seizures are more frequent during the CVT comparing to the other types of stroke. Seizures occur mainly in cases of parenchymal lesions, SSS thrombosis and in patients having motor or sensitive deficit [5].

Most of the routine blood assessment does not have diagnostic value for CVT. However, they may help establish its etiology. Several studies confirmed the elevation of the D-dimers during the CVT but their predictive negative value seems only interesting for patients having signs of encephalic reach. Therefore, an absence of D-dimers elevation in case of isolated headache can not rule out CVT [6].

In our study, the cerebrospinal fluid (CSF) mostly showed abnormal composition and elevation of the intracranial pressure. The CSF assessment revealed associated purulent meningitis in two patients, meningitis with an increased number of lymphocytes and red cells, an isolated elevation of red cells at two patients. However, we did not find any elevation of proteins in CSF. The opening pressure was elevated in all patients except the cases of cavernous sinus thrombosis. It varies between 19 and 48 cm H2O.

Detailed CT scan findings have been described in earlier literature [7-10]. The CT scan without and with of contrast agent injection remains the first exam achieved whenever a CVT is suspected. It provides the diagnosis proof and allows showing the direct signs of CVT. However, up to 20% CVT cases have shown a normal CT [11], more frequently they are patients with isolated intracranial hypertension. Therefore, a CVT diagnosis has to be systematically discarded before retaining an idiopathic intracranial hypertension diagnosis [12, 13]. However, the CT scan sensitivity was 30% in Ameri study [14]. The empty delta sign (the frequent direct sign) was found in six patients (23%) compared to 20% found in the literature [15]. The empty lateral sinus sign is the equivalent of the empty delta sign. This seemed to be underestimated in the literature [16], whereas it represents the first direct sign in our study and was found in 11 patients (37%). It is necessary to differentiate it from hypoplastic sinus mainly the left lateral sinus. A special attention must be allocated to the signs of the transverse sinus on the mastoid in images before contrast agent injection.

Indirect signs are more frequent and less specific compared to direct signs in different studies. This was also confirmed in our study (85%). Most often, the CT-scan shows thrombosis consequences on the cerebral parenchyma under the shape of a hypodensity (oedema or venous infarction), or a hyperdensity bound to a haemorrhage going from some small patches to a real intraparenchymal haematoma. The infarct is frequently hemorrhagic (10 to 50% of cases in the literature) [17]. In our study, the CT scan was sufficient to diagnose 50% of cases at the admission.

Currently, the gold standard in the diagnosis of CVT is to combine MRI and MRV for visualizing the thrombosed vessel. The indirect signs of CVT shown in the MRI are similar to those in the CT scan since they are not specific; both might withdraw 25% of cases. The sensitivity of the MRI associating the MRV is 90% in Lafitte study [18]. The MRI and the MRV allowed the diagnosis in all cases of our series (56%). The MRI alone is limited by flow artifacts that can lead to false positives and the absence of hyperintense signal on T1 and T2-weighted images at the onset of acute thrombosis [19]. During the first 3 to 5 days the thrombosed sinus is isointense on T1 and hypointense on T2, and thus very difficult to differentiate from normal veins. All MRI examinations have been achieved after the 5th day of the onset. For all cases; (except those with deep veins thrombosis, cortical veins thrombosis and the cavernous sinus thrombosis); the thrombosed sinus was hyperintense on T1 and hyper or isoointense on T2. MRV done alone cannot discriminate hypoplasia than lateral sinus thrombosis [20]. It is illustrated by one case (case number 5) that shows an amputation of the hypoplasic LLS confirmed by morphological T2 and T1 sequences. The cortical or deep vein thrombosis might be misdiagnosed even by MRI and MRV. This fact occurred in one case presented with a deep vein thrombosis (case number 18).
The diagnosis of isolated cortical vein thrombosis might be difficult, requiring conventional angiography. In our series, conventional angiography investigation was not used. The Echo-planar susceptibility-weighted images (T2*) are particularly useful in isolated cortical venous thrombosis and during early stage of acute CVT when T1 and T2 lack sensitivity [21]. The diagnosis of cortical venous thrombosis was performed by echo-planar susceptibility-weighted images (T2*) at one patient (case number 20) (Figure 4). The thrombosis of the cavernous sinus was revealed in three cases by an absence of heightening of one or the two cavernous sinuses in the coronal T1 images after Gadolinium injection, it was the best to show the sinus thrombosis. Two patients had cavernous sinus thrombosis associated with intracavernous carotid aneurysm.

CVT is often multifactorial since several disorders might initiate or dispose patients. In the International studies of the Cerebral Vein and Dural Sinus Thrombosis (ISCVT), 44% of the patients had shown more than one origin and predisposing factor including the congenital and genetic thrombophilia which was recorded in 22% patients [22]. In developed countries, non-infectious origins such oral contraceptives, cancer, and thrombophilia are the most frequent causes with a prevalence of 80% [4, 23]. The etiologies in our study was similar to the developing countries findings [24, 25]. Infectious and post-partum causes are the most frequent causes. Local infectious etiologies are observed in seven cases including four cases of otomastoiditis, three cases of severe staphyllococcal of the face and one case of ethmoiditis. CVT in these cases may be explained by the consultation delay of our patients. CVT represents 23.5% of the stroke during pregnancy and postpartum [26]. Several factors could be associated to the postpartum CVT, these included caesarean section, arterial hypertension, and anemia. On the other hand, eclampsia is not a risk factor. Indeed no case of eclampsia has been counted in our study [27]. In one case, CVT occurred after epidural analgesia. Oral contraceptives represent 12% in Biousse and Bousser study. In our context, we are convinced that some cases are not diagnosed considering the increasing use of the oral contraceptives in our environment. Hemostasis investigation includes congenital or genetic thrombophilia (deficiencies in antithrombin III, protein C, and protein S and having the factor V Leiden or prothrombin gene mutations, and antiphospholipid antibodies). We have investigated thrombophilia in four cases; one case of deficiency in protein C, and two cases in protein S were revealed. The protein S deficiency was not confirmed by a second dosage. Congenital thrombophilia is certainly underestimated in our study. Indeed, due to Behcet disease is frequent in Mediterranean countries, but Behcet's cases seem less frequent in our study compared with Daif study. The low frequency of Behcet disease might be explained by our limited cases. CVT was idiopath in 20% in agreement with literature data [28].

Heparin (low-molecular-weight heparin or standard heparin) is the first-line treatment even in presence of hemorrhagic infarction [29, 30]. In our study, 90% of patients benefited from anticoagulant treatment including those with hemorrhagic infarction. Systemic or local thrombolysis may be used in CVT when patients deteriorate despite adequate anticoagulation and other causes of deterioration have been ruled out, thrombolysis or thrombectomy may be considered in specialized centers [31, 32]. In patients with isolated intracranial hypertension, a lumbar puncture for removing the cerebrospinal fluid is required before starting heparin. If intracranial pressure is severely raised, the general recommendations should be followed, this starts with manitol treatment to admission to an Intensive Care Unit with intracranial pressure monitoring or even decompressive hemicraniectomy when a risk of cerebral herniation exists. In our patients who consulted at a stage of important decrease of visual cut, y we used carbonic anhydrase inhibitor or manitol or even steroids although they are not recommended by most authors [33]. Antiepileptics and antibiotics are useful to treat seizures and infection. One of our patient underwent a partial epilepsy which was stabilized under carbamazepin. The advisable anticoagulant treatment duration is 6 to 12 months. For our patients, the mean duration of anticoagulation therapy was 6 months. ISCVT shows that death and severe sequelae were observed in 15% of patients [22]. In our study, the result was similar to ISCVT (13% vs 15%) with two cases of severe sequelae and two cases of death.

**Conclusion**

CVT is not uncommon in Morocco. Post-partum and infectious causes are more frequent in this study. Behcet's disease etiology seems to be underestimated. The investigation of the haemostasis should be a routine after each cerebral venous thrombosis. MRI and Venous MR angiography are the key diagnostic tools. CT scan should be more effective for establishing the CVT diagnosis. The outcome is favorable using heparin treatment.

**Competing interests**

The authors declare no competing interests.

**Authors’ contributions**

Souiri Z is the chief investigator and manager of the study. He drafted this manuscript. Other authors are co-investigators. All authors read and approved the final version of the manuscript.

**Tables and figures**

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Table 1: Patients data

| N  | Age, gender | Clinical symptoms | Causes and risk factors | Topography | Parenchyma involvement | Outcome |
|----|-------------|------------------|------------------------|------------|-----------------------|---------|
| 1  | 11, M       | ICH, P, IIIrd np, VIth np, exophtalmos, | Severe staphylococcal of the face | SC         | no                    | normal  |
| 2  | 27, F       | ICH, P, motor deficit, DC | Post-partum, otomastoiditis | SSS, SLD   | yes                   | Motor deficit |
| 3  | 60, M       | Seizures, motor deficit, DC, fever | absent | SLD         | yes                   | normal  |
| 4  | 24, F       | Ptosis, headache, IIIrd np, VIth np, exophtalmos, DC, fever | Severe staphylococcal of the face | SC         | no                    | IIIrd np, VIth np |
| 5  | 17, F       | ICH, P, VIth np, exophtalmos, fever | Probable local infection | SC         | no                    | death   |
| 6  | 27, F       | ICH, P, seizures, motor deficit, | Post-partum | SSS         | yes                   | seizures |
| 7  | 43, M       | Seizures, motor deficit, | Absent | SSS, SLG    | yes                   | normal  |
| 8  | 28, F       | ICH, seizures, motor deficit, DC | Post-partum | SSS         | yes                   | death   |
| 9  | 21, F       | ICH, fever | Post-partum | SSS         | no                    | normal  |
| 10 | 17, M       | ICH, P, | Behçet disease | SLD         | No                    | normal  |
| 11 | 18, F       | Headache, status epilepticus, motor deficit, DC, fever | Post-partum | SSS, SLD    | Yes                   | normal  |
| 12 | 18, F       | ICH, P, fever | Otomastoiditis | SLG         | No                    | normal  |
| 13 | 46, M       | ICH, IIIrd np, VIth np | Severe staphylococcal of the face | SC         | No                    | Blindness, IIIrd np, VIth np |
| 14 | 54, M       | ICH, P, motor deficit | Pulmonary carcinoma with metastasis | SLD         | yes                   | death   |
| 15 | 25, F       | ICH, P, status epilepticus, DC, fever | Post-partum | SSS, SLD    | yes                   | normal  |
| 16 | 24, M       | Headache, fever, IIIrd np, VIth np exophtalmos | Ethmoiditis | SC         | no                    | normal  |
| 17 | 65, F       | DC | Miliary tuberculosis | SLD         | no                    | normal  |
| 18 | 25, F       | Status epilepticus, DC, fever | absent | Basilar veins, ICV | yes | Cognitive dysfunction, motor deficit |
| 19 | 13, M       | ICH, P | otomastoiditis | SLD         | No                    | normal  |
| 20 | 40, F       | Headache, motor deficit | Post-partum, epidural analgésia | Cortical vein | No | normal |
| 21 | 7, F        | ICH, P, fever | Absent | SSS, SLD, SLG | No | normal |
| 22 | 55          | ICH, motor deficit, DC | Absent | SLG         | Yes                   | death   |
| 23 | 18 months, F | Status epilepticus, hyporeactivity | Protein C deficiency | SSS         | yes                   | Pyramidal hypertonia |
| 24 | 40, F       | Headache, seizures, motor deficit | Post-partum | SSS         | yes                   | Motor deficit |
| 25 | 19, F       | Headache, DC | Post-partum, | SLD         | No                    | normal  |
| 26 | 18, F       | ICH, P | Essential thrombocytémia | SSS         | No                    | normal  |
| 27 | 44, F       | ICH, motor deficit, fever | absent | SSS, RLS    | no                    | normal  |
| 28 | 24, F       | Headache, DC | Post-partum | SSS, LLS    | yes                   | normal  |
| 29 | 28, M       | Motor deficit | Behçet disease | Basilar veins | yes | Motor deficit |
| 30 | 23, F       | ICH, seizures, DC, fever | absent | SSS         | yes                   | normal  |

IIIrd np, third nerve paralysis; DC, disorder of consciousness; ICH, intracranial hypertension; ICV, internal cerebral veins; P, papilledema SSS, superior sagittal sinus; LLS, left lateral sinus; RLS, right lateral sinus; CS, cavernous sinus; F: feminine; M: masculine.
### Table 2: The 14 patients with both CT scan, MRI and MRV

| Case | CT scan                                      | MRI                                           | MRV                  | Thrombosed sinus or vein |
|------|----------------------------------------------|-----------------------------------------------|----------------------|--------------------------|
| 1    | Absence of bilateral CS enhancement,         | Absence of bilateral CS enhancement,         | normal               | CS                       |
|      |                                              | aneurysm of left intracavernous carotid       |                      |                          |
| 3    | Cortical and subcortical hypodensity         | HS T1, T2 RLS; venous infarction              | Amputation RLS       | RLS                      |
| 5    | normal                                       | Absence of bilateral CS enhancement,         | Amputation LLS       | CS                       |
|      |                                              | venous infarction.                            | (hypoplasia).         |                          |
| 6    | Cortical and subcortical hypodensity,        | Iso T1, hypo T2 SSS; hemorrhagic infarction   | Amputation SSS       | SSS                      |
|      | multiple grouped hyperdensities              |                                               |                      |                          |
| 14   | Empty lateral sinus sign,                    | HS T1, T2 RLS; multiple metastasis            | Amputation RLS       | RLS                      |
|      | enhancement of sinus wall, cortical          |                                               |                      |                          |
|      | and subcortical hypodensity                  |                                               |                      |                          |
| 16   | Absence of bilateral CS enhancement,         | Absence of right CS enhancement;             | Normal               | CS                       |
|      | enhancement of CS wall                       | aneurysm of right intracavernous carotid      |                      |                          |
| 17   | Cord sign, empty right lateral sinus sign.   | HS T1, T2 RLS; multiple tuberculoma.          | Amputation RLS       | RLS                      |
| 18   | bilateral venous infarction (Rosenthal vein  | Partial obstruction of LLS (T1 Gadolinium),   | Normal               | Rosenthal vein and      |
|      | and intern cerebral vein).                   | bilateral venous infarction (Rosenthal vein   |                      | intern cerebral vein     |
|      |                                              | and intern cerebral vein).                    |                      |                          |
| 20   | normal                                       | HS FLAIR, hyposignal T2* of a cortical vein.  | normal               | Cortical vein            |
| 21   | Empty delta sign, empty lateral sinus sign   | HS T1, T2 RLS, LLS, SSS                       | Amputation SSS,      | SSS, RLS, LLS            |
|      | (right and left)                             |                                               | RLS, LLS             |                          |
| 24   | Hematoma                                     | Hematoma                                      | Defects SSS          | SSS                      |
| 25   | Empty right lateral sinus sign               | HS T1, T2 RLS                                | normal               | SSS                      |
| 26   | normal                                       | HS T1, isosignal T2 SSS                       |                      |                          |
| 28   | Cortical and subcortical hypodensity,        | hemorrhagic infarction                        | Defects SSS, LLS     | SSS, LLS                 |
|      | multiple grouped hyperdensities              |                                               |                      |                          |

HS, hypersignal intensity; SSS, superior sagittal sinus; LLS, left lateral sinus; RLS, right lateral sinus; CS, cavernous sinus

**Figure 1:** (case 21) Sagittal MR T1 sequences without Gadolinium injection showing: hypersignal of thrombosed SSS on the right (black arrows) (A), hypersignal of thrombosed LLS on the left (white arrow) (B)
Figure 2: Principal clinical signs of the 30 patients

Figure 3: (case 14) CT scan after contrast product injection, axial sequence of the posterior fossa showing empty right lateral sinus (arrow)
Figure 4: (case 20): FLAIR MR cerebral axial sequences (on the left) showing a hypersignal at the level of the ascending frontal gyrus (arrow) and in T2* (on the right) showing a hyposignal of a cortical vein (arrow) which is thrombosed.

Figure 5: Etiologies of the 30 patients