CASE REPORT

An Outpatient Approach to the Management of Recurrent Epistaxis in Patients with Hereditary Hemorrhagic Telangiectasia

Vasilios Chalkiadakis, Silvestros Boukouvalas, Maria Manitsopoulou, Konstantinos Papadopoulos, Georgios Karatzias

ABSTRACT

Patients with Rendu–Osler–Weber syndrome, also known as HHT, have a prevalence of approximately 1 in 5,000 to 10,000, with a large geographic variability. The syndrome is inherited in an autosomal dominant manner with incomplete penetrance, although in 20% of cases, there is no family history. Many genes have been implicated with the syndrome, but the ENG in chromosome 9 seems to be the most significant.

Even though genetic tests are available, diagnosis is based on the Curacao criteria, established by the Scientific Division of the Hereditary Hemorrhagic Telangiectasia International Foundation. This condition is characterized by multiple telangiectases, spontaneous and recurrent epistaxis, visceral lesions, and the presence of a first-degree relative with Rendu–Osler–Weber syndrome. The diagnosis is considered definite when three or more findings are present; possible or suspected when two findings are present; and finally, unlikely, when only one or none of these findings are present. The positive predictive value for a definite clinical diagnosis is 100%, thus genetic tests are more essential in individuals with a possible clinical diagnosis of the syndrome. Clinically, telangiectases usually appear mucocutaneous with a very characteristic development. These often present in the nasal and gastric mucosa, on the lips, tongue, and fingers. Moreover, most patients exhibit larger arteriovenous malformations located in the hepatic, pulmonary, or cerebral vasculature.

Bleeding complications may arise from each arteriovenous malformation or telangiectasia. Spontaneous and recurrent epistaxis is the most common symptom in these patients. It affects up to 98% and most of them will have their first episode of nasal bleeding before the age of 20. Usually, these episodes are minimally bothersome, but in some cases, the severity might increase by time. Mucosal telangiectases have usually a cosmetic concern, whereas those in the gastrointestinal tract might cause chronic bleeding and iron-deficiency anemia. Pulmonary, hepatic, and cerebral vascular malformations are more important in terms of morbidity and mortality, although these are less common in individuals. Even though Rendu–Osler–Weber is a multisystemic syndrome, otorhinolaryngologists may frequently be involved in order to control nasal bleeding.

There are multiple different options for the treatment of epistaxis in these patients. Most of the management decisions focus on bleeding complications. In the acute setting, clinicians usually proceed to anterior or posterior nasal packing with absorbable or nonabsorbable materials. Other options include conservative therapy, such as saline rinses, barrier creams, and topical emollients.
Those therapies are considered as noninvasive, because the risk for the patient is minimal. Furthermore, medical therapies can be applied, such as antiestrogen drugs, bevacizumab, or timolol.\textsuperscript{7-9} When the bleeding episodes become more severe, even life-threatening, surgical options are considered. These procedural approaches include coagulation, coblation, septodermoplasty, embolization, or even nasal closure, known as the Young’s procedure.\textsuperscript{10-12} These treatments aim at reducing the severity and the frequency of bleeding episodes and none of them has obtained full success. Furthermore, all these options come up with various risks for the patients, so a stepwise approach should be considered.

Within the last year, three patients previously diagnosed with HHT were referred in our clinic for recurrent episodes of nasal bleeding. The aim of this study is to describe our strategy for the management of epistaxis without the need of hospitalization. Our protocol is based upon coblation via RF energy, and in contrast to other medical centers, it is performed in an outpatient setting under local anesthesia, without intravenous sedation.

\textbf{CASE REPORT}

In our attempt to get a detailed picture of each patient, a complete medical history is followed by a thorough examination in order to underline all possible manifestation sites of the syndrome. According to the clinical findings and prior screening, blood testing, imaging, and echocardiogram are further recommended, in order to ensure that all severe arteriovenous malformations are found. Patients with suspected gastrointestinal bleeding are advised to take a gastrointestinal endoscopic investigation.

The method that we performed included firstly chemical coagulation using nitric acid solution under local anesthesia in order to locate the accurate site of telangiectasia in the nasal cavity. The subsequent stage was coblation via RF energy, which breaks down molecular bonds at low temperatures (70°C). Telangiectasia cases were managed due to a RF generator with nonstick forceps in bipolar coagulation mode, at the lowest possible energy setting.

Patients were admitted for treatment sessions every 10 days. In general, three to four sessions were necessary for the complete management of each case. The patients were also encouraged through training courses to use on their own a local moisturizing and hemostatic ointment every day, for regular nasal mucosal care.

As mentioned earlier, three patients diagnosed with Rendu–Osler–Weber syndrome were recently treated by coblation via RF energy, for recurrent epistaxis. As noted in Figure 1, they all belong to the same family. Two of them are first-degree relatives, while the third one is a second-degree relative with one and a third-degree relative with the other. Until now, these three patients had no other treatment, except nasal packing and local ointments in order to control acute bleeding from the nose.

A 45-year-old female presented to our clinic with lifelong complications associated with Rendu–Osler–Weber syndrome. Her medical history revealed multiple telangiectases in the nasal mucosa, recurrent episodes of severe epistaxis, which occurred daily for a 2-month period and a family history of HHT. Furthermore, this chronic nasal bleeding contributed to iron-deficiency anemia, a common condition in these cases. No arteriovenous malformations were identified in the pulmonary, hepatic, or cerebral vasculature and gastrointestinal tract. For the management of these bleeding episodes, three treatment sessions were performed.

The second case is the 71-year-old mother of the individual described above. This patient presented with three Curacao criteria, including arteriovenous malformations in the gastrointestinal tract, classic telangiectases in the nasal cavity, and recurrent epistaxis. On physical examination, the patient reported systemic hypertension on medication and a previous episode of gastrointestinal bleeding due to a telangiectasia in duodenum. The patient referred several episodes of epistaxis, which were increased in frequency to 2 or 3 per month in the last 3 years. Four sessions were performed, in one of which, nasal packing for 48 hours was required.

Finally, the third patient presented with possible diagnosis of HHT. He experienced recurrent episodes of epistaxis in the last few years, associated with physical activity, as well as multiple nasal and oral mucosal telangiectasias. Interestingly, this 40-year-old patient is nephew of the last patient mentioned (second-degree relative), and therefore first cousin of the primary case described (third-degree relative). The management
strategy included three sessions, whereas nasal packing was also implemented once for 48 hours.

DISCUSSION

Hereditary hemorrhagic telangiectasia, also known as Rendu–Osler–Weber syndrome, is a relatively common systemic fibrovascular dysplasia, inherited in an autosomal dominant manner with incomplete penetrance. The major genes that have been identified are: ENG encoding endoglin (HHT type I), ACVRL 1 encoding activin receptor-like kinase (ALK-1) (HHT type II) and SMAD 4 encoding Smad 4 (HHT in association with juvenile polyposis). It is considered that over 80% of these patients will have mutations in the ENG or ACVRL 1 gene. This condition occurs in heterozygotes, while the homozygote form is not compatible with life.

The mutated genes encode membrane glycoproteins which compose the transforming growth factor beta. All these functions are essential in angiogenesis. As a result, there is proliferation of endothelial cells, excessive vessel branching, and decreased recruitment of mural cells. This leads to the formation of telangiectasia, a systemic fibrovascular dysplasia. All these malformed vessels extend through the dermis and have excessive layers of muscle without elastic fibers, which is the basic defect in these patients. Due to this abnormal wall structure, these vessels are prone to spontaneous ruptures and injuries.

Even though genetic tests are nowadays available, diagnosis remains clinical, based on the Curacao criteria. These criteria include not only the classic triad of recurrent and spontaneous epistaxis, multiple telangiectasates at characteristic sites, and family history with an affected first-degree relative, but also the presence of large arteriovenous malformations, found in the gastrointestinal tract, or in the pulmonary, hepatic, spinal, and cerebral vasculature. The diagnosis is confirmed when at least three of these findings are present. Due to these important lesions in terms of mortality and morbidity, current guidelines recommend careful screening and close follow-up monitoring of all patients.

These individuals are at high risk for bleeding at the site of telangiectases or visceral lesions. The most frequent form of bleeding is epistaxis, affecting almost 100% of the patients. These nasal bleeding episodes are recurrent and the intensity of this symptom may range from minimally bothersome to life-threatening. Furthermore, in some cases, this chronic bleeding may lead to iron deficiency anemia, or even multiple blood transfusions may be required.

Nowadays, treatment is only supportive and the aim is to prevent any complications. Moreover, there is still no consensus for the best option. These options vary from conservative, medical, or surgical/procedural. Usually, majority of clinicians proceed to anterior-posterior nasal packing or cauterezation when a bleeding episode occurs, although others prefer to manage the disease for as long as possible. In more severe cases, definitive treatment options are performed. These options include arterial embolization, laser, septodermoplasty, and nasal closure (Young’s procedure). Because of the serious side effects, these options stand as the last resort that clinicians implement. Our treatment protocol is based on prevention of epistaxis through extensive information of patients, training courses for self-care, and also on a procedural control of mucosal telangiectases by coblation via RF energy.

In contrast to other medical centers, we perform this therapy in an outpatient setting, under local anesthesia without intravenous sedation. One significant benefit is that patients do not undergo the risk of recurrent general anesthesia, while this procedure is also time-effective for the individuals, as they are not hospitalized. Although the use of local anesthesia minimizes the pain level, there is a possibility for some patients to mention sensitivity at the site of intervention.

Another interesting advantage of this procedure is the lack of nasal occlusion by tape or packing. Even a non-permanent closure of the nostrils could cause xerostomia, anosmia, sneezing, mucosal infection, and formation of crusts. Nevertheless, there is always the risk of rupturing arteriovenous shunts in the nasal mucosa during application or removal. Our personal experience showed that patients were very satisfied with this adjustment.

Furthermore, due to RF energy, coblation can be achieved at relatively low temperatures (70°C), thus the risk of damaging the surrounding, healthy mucosa is minimized. In addition to this, thermal injury is decreased in the deeper layers of nasal mucosa, which leads to less crust formation. Besides, no relative scarring or septal perforation has been reported to date in our patients by the implementation of this regular therapy. Finally, follow-up of these three patients has shown that both the frequency and the severity of bleeding episodes have been reduced. Within the last 6 months, none of them has mentioned a new episode of epistaxis, while they all reported that with our approach, their quality of life has been improved.

CONCLUSION

Rendu–Osler–Weber syndrome is a multisystemic disease, characterized by the presence of vascular malformations. As a result, spontaneous and recurrent epistaxis is often reported in these patients. This may range from minimal bothering to life-threatening bleeding episodes. The majority of management decisions concern local complications caused by these episodes. Several procedural options are available for the treatment of nasal bleeding. Thus, it is critical that clinicians should be up-to-date in
An Outpatient Approach to the Management of Recurrent Epistaxis in Patients


REFERENCES