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Giant Right Liver Hemangioma associated with Kasabach-
Merritt Syndrome in an Adult Patient

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## **Case Study**

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7 **Introduction:** Liver hemangiomas are often asymptomatic and diagnosed incidentally.

- 8 Kasabach-Merritt syndrome (KMS) or consumptive coagulopathy is a rare but life-threatening
- 9 complication of liver hemangioma occurring during observation. Surgery is an appropriate
- treatment option in such condition and coagulation usually returns to normal after surgical
- excision. We herein report a case of giant right liver hemangioma with Kasabach-Merritt
- syndrome treated surgically with literature review.
- 13 Case Presentation: A 36 –year old woman with a giant liver hemangioma (20 cm) discovered
- three years ago, who presented to emergency department for pallor and fatigability and no
- abnormalities were found on physical examination. After excluding hematologic diseases, a
- 16 Kasabach-Merritt syndrome associated with giant liver hemangioma had been
- 17 retained.Csoagulation disorders returned to normal after successful surgical resection of lesion
- by performing a right hepatectomy.
- 19 **Conclusion:** Resection is an appropriate and effective surgical procedure to treat giant liver
- 20 hemangioma associated with Kasabach-Merritt syndrome.
- 21 **Keywords:** giant liver hemangioma consumptive coagulopathy, surgical resection

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#### Introduction

- Hemangiomas are the most common benign tumor of liver. Most liver hemangiomas are
- asymptomatic, small (< 4cm) and diagnosed incidentally [1]. A liver hemangioma is qualified
- 27 giant when it has a diameter greater than 10 cm. Asymptomatic liver hemangioma is managed
- 28 conservatively. However symptomatic or complicated lesion justified the surgical management
- 29 [2,3]. Kasabach-Merritt syndrome is a rare complication of liver hemangioma and it presents
- 30 as hemolytic anemia, thrombocytopenia, prolonged prothrombin time, and
- 31 hypofibrinogenemia. Surgical treatment is an appropriate therapeutic option for such condition
- and coagulation usually returns to normal after surgical removal . We report a case of giant
- 33 right liver hemangioma associated with Kasabach-Merritt syndrome treated surgically with
- 34 literature review.

### **Case Presentation**

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- A 36 –year old woman presented to emergency department for pallor and fatigability. A giant
- 38 liver hemangioma was discovered incidentally during pregnancy three years ago that it was
- 39 managed conservatively. At admission, the patient was pale but not icteric and no abnormalities
- were found on the physical examination. Laboratory testing revealed that blood count and liver
- 41 function tests results were WBC:  $2.99.\ 10^9 / L (4.0-10.010^9 / L)$ , Hemoglobin: 8.2 g/L (115-150 m)
- 42 g/L), Platelets:  $80.000/\text{mm}^3$ (110–320.) ALT: 18 m/L (0–40 m/L), AST: 21 m/L (0–42
- 43 m/L), ALP: 56 m/L (40–150), GGT: 35 m/L (0–52 m/L), TB: 9.7mmol/L (5.0–
- 44 21.0mmol/L), DB: 4.8mmol/L(0.0-7.0mmol/L), Fibrinogen: 1.83, g/L (2.00-4.00, g/L,
- 45 INR: 1.54 (0.85–1.50), Prothrombin time: 18,2 sec (11–15). Hepatitis B virus and hepatitis
- 46 C virus markers were negative, and  $\alpha$ -fetoprotein level was 8 ng/dL( 0–10 ng/dL).
- 47 As showed on Computed Tomography Scan, the lesion occupied almost all liver segments 5, 6,
- 48 7 and 8 and measuring approximately 20 x 12 x 8 cm without vessel compression (Fig. 1)
- 49 .After excluding hematologic diseases such as hemolytic anemia, hemolytic uremic syndrome,
- 50 , systemic inflammatory response syndrome and basing on laboratory results, a Kasabach-
- 51 Merritt syndrome associated with giant liver hemangioma had been retained.
- The hematologic abnormalities had been corrected before surgery by using packed red blood
- 53 cell, platelet concentrate and fresh frozen plasma. The operative exploration found a huge
- reddish pink tumor with thin walls and occupying almost all the right hemiliver. The intra-
- operative decision was to perform a right hepatectomy. The right liver portal vein ant artery
- 56 was clamped after liver helium dissection (fig.2) and parenchymal transection was performed
- 57 using an ultrasonic dissection device. The tumor was dissected away from the the inferior vena
- 58 cava (IVC) after exposure of the antero-medial surface of the IVC and ligation of several short
- 59 hepatic veins. The right hepatic vein, the right portal vein and artery were the last vascular
- 60 elements to be divided. The tumor had a length of 20 cm approximately (fig.3). The patient
- 61 developed a right bloody pleural effusion which was resolved after thoracic drainage
- 62 maintained during five days. Histological examination of operative specimen revealed a
- 63 cavernous hemangioma. The coagulation and hematologic abnormalities returned to normal
- value 3 weeks after surgery (Table.1).

#### Discussion

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- Hemangiomas are one of the most common benign tumors of liver. According to their size,
- 69 hepatic hemangiomas are classified into 3 types: small (<5 cm), large (5–10 cm), and giant

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       (>10 cm). Observation is justified in asymptomatic lesion and surgery is indicated in the
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       presence of complications [4]. Consumptive coagulopathy or Kasabach-Merritt syndrome
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       (KMS), described firstly by Kasabach and Merritt in 1940, is a rare and severe coagulation
       disorder associated with vascular malformations [5]. The Kasabach-Merritt syndrome is
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       characterized by thrombocytopenia, hemolytic anemia, and consumptive coagulopathy [6].
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       Surgical management remains an effective and curative treatment for complicated or
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       symptomatic liver hemangioma [7]. Our patient underwent a right hepatic liver resection using
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       a hanging manoeuvre to avoid difficulties and minimize risk of bleeding during liver
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       mobilisation. Transfusion of three units of red blood cells was required because of preexistent
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       anemia and operative blood loss (300 ml). Risk of operative bleeding is likely to be more
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       related to hemangioma size (> 20 cm) [7]. Compression of major vessels surrounding the
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       lesion may expose to high risk of uncontrolled severe bleeding and blood loss during operation.
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       So, cell saver system is highly recommended to decrease blood transfusion rate in these
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       patients. The liver resection procedure is more likely recommended to remove liver
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       hemangioma associated with KMS because hemangioma often has an extremely greater size
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       (>20cm) making liver mobilisation more difficult with high risk of bleeding. So preligation of
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       both artery and portal vein decreases lesion size, facilitates liver mobilisation and thus
       reduces risk of bleeding. In addition, an extremely giant hemangioma can occupy entirely a
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       hemiliver or more and performing anatomic liver resection will not lead to substantial loss of
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       healthy liver parenchyma. Although surgery remains the radical treatment of liver
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       hemangioma, other therapeutic options including transcatheter arterial embolization (TAE)
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       and radiofrequency ablation can especially be considered in patients with high surgical risk
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       [8-10]. These therapies can be performed prior to surgery in order to reduce tumor size of
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       extremely giant lesion [8-10]. As reported, liver transplantation had the same effects as
       surgery in the treatment of Kasabach- Merritt syndrome associated with liver hemangioma
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       [11,12]. However, liver donor is rare, and patient needs to take an immunosuppressive
       treatment for a long-term period after transplantation. Since 2008, oral propanolol have been
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       used to treat hepatic hemangioma and it was largely used in combination with steroids in
       infant [13,14]. As demonstrated by published study results, the efficacy and safety of this
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       therapy as a first treatment, had an positive impact on changing the classical therapeutic
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       indications particularly in diffuse hepatic lesion (type 3) by obviating liver transplantation for
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       many of these patients [14,15], therefore, the usage of propanolol alone or in combination
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       with steroids as first treatment line in infantile liver hemangioma resulted in decreasing the
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       indications of surgical treatment options. On other hand and in adult patient, propranolol has
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- been used as a first therapeutic option for liver hemangioma in selective few patients with
- good results. However randomized prospective studies are highly recommended to evaluate
- the results and clarify the appropriate use of this agent in such condition [16,17]
- 107 Conclusion
- In summary ,Kasabach-Merritt syndrome is an uncommon complication of liver hemangioma
- occurred in adult patient. Surgery is an effective therapeutic option and hematological
- abnormalities and coagulation disorders returned to normal values after surgical resection. In
- such condition; liver anatomic resection is a safer surgical procedure.
- 112 Disclaimer regarding Consent/Ethical Approval:
- As per university standard guideline participant consent and ethical approval has been
- collected and preserved by the authors.
- [ 13] Aly MM, Hamza AF, Abdel Kader HM, Saafan H, Ghazy MS, Ragabl A.
- Therapeutic superiority of combined propranolol with short steroids course over
- propranolol monotherapy in infantile hemangioma. Eur J Pediatr.
- 118 2015;174(11):1503-1509.

- [14] Shah SD, Baselga E, McCuaig C, et al. Rebound growth of infantile
- hemangiomas after propranolol therapy. Pediatrics. 2016;137(4).doi:
- 122 10.1542/peds.2015-1754.

123

- [15] Sarıalioğlu F, Erbay A, Demir S. Response of infantile hepatic hemangioma to
- propranolol resistant to high-dose methylprednisolone and interferon-α
- therapy. Pediatr Blood Cancer. 2010;55(7):1433-1434.

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#### 129 References

- [1] Choi BY, Nguyen MH. The diagnosis andmanagement of benign hepatic tumors. J
- 132 ClinGastroenterol 2005;39:401–12.
- [4]. Hoekstra LT, Bieze M, Erdogan D, Roelofs JJTH, Beuers UHW, van Gulik TM.
- Management of giant liver hemangiomas: an update. Expert Rev GastroenterolHepatol.
- 135 2013;7:263.
- 136 [3]. Toro A, Mahfouz A-E, Ardiri A, et al. What is changing in indications and treatment of
- hepatic hemangiomas. A review. Ann Hepatol. 2014;13:327.
- 138 [4]. Weimann A, Ringe B, Klempnauer J, Lamesch P, Gratz KF, Prokop M, Maschek H, Tusch
- G, Pichlmayr R: Benign liver tumors: differential diagnosis and indications for surgery. World
- 140 J Surg1997, 21:983-990.
- 141 [5]. Kasabach HH, Merritt KK. Capillary hemangioma with extensive purpura. Am J Dis
- 142 1940;59:1063-1070.

- [6]. Aslan A, Meyer ZuVilsendorf A, Kleine M, Bredt M, Bektas H. Adult Kasabach-merritt
- syndrome due to hepatic giant hemangioma. Case Rep Gastroenterol 2009;3:306-312.
- [7]. Oak CY, Jun CH, Cho EA, et al. Hepatic hemangioma with Kasabach-Merritt syndrome
- in an adult patient. Korean J Gastroenterol 2016;67:220–3.
- [8]. Seo HI, Jo HJ, Sim MS et al. Right trisegmentectomy with thoracoabdominal approach
- after transarterial embolization for giant hepatic hemangioma. World J
- 149 Gastroenterol.2009;15:3437–3439
- 150 [9]. Zhou J-X, Huang J-W, Wu H, Zeng Y. Successful liver resection in a giant hemangioma
- with intestinal obstruction after embolization. World J Gastroenterol. 2013;19:2974.
- 152 [10]. Gao J, Ke S, Ding X, Zhou Y, Qian X, Sun W. Radiofrequency ablation for large hepatic
- hemangiomas: initial experience and lessons. Surgery. 2013;153:78.
- 154 [11]. Hochwald SN, Blumgart LH. Giant hepatic hemangioma with Kasabach-Merritt
- syndrome: is the appropriate treatment enucleation or liver transplantation? HPB Surg 2000;
- 156 11:413-419.
- 157 [12]. Meguro M, Soejima Y, Taketomi A, et al. Living donor liver transplantation in a patient
- with giant hepatic hemangioma complicated by Kasabach-Merritt syndrome: report of a case.
- 159 Surg Today 2008;38:463-468.
- 160 [13] Aly MM, Hamza AF, Abdel Kader HM, Saafan H, Ghazy MS, Ragabl A. Therapeutic
- superiority of combined propranolol with short steroids course over propranolol monotherapy
- in infantile hemangioma. Eur J Pediatr. 2015;174(11):1503-1509.
- 163 [14]Shah SD, Baselga E, McCuaig C, et al. Rebound growth of infantile hemangiomas after
- propranolol therapy. Pediatrics. 2016;137(4).doi: 10.1542/peds.2015-1754.
- 165 [15] Sarialioğlu F, Erbay A, Demir S. Response of infantile hepatic hemangioma to propranolol
- resistant to high-dose methylprednisolone and interferon-α therapy. Pediatr Blood Cancer.
- 167 2010;55(7):1433-1434.
- [16] Mazereeuw-Hautier J, Hoeger PH, Benlahrech S et al. Efficacy of propranolol in hepatic
- infantile hemangiomas with diffuse neonatal hemangiomatosis. J Pediatr 2010; 157:340–342.
- 170 [17] Amal Mhanna, Wayne H. Franklin, Anthony J. Mancini. Hepatic Infantile Hemangiomas
- 171 Treated with Oral Propranolol—A Case Series. Pediatric Dermatology Vol. 28 No. 1 39–45, 2011
- 173174

# Table 1: per and postoperative results of blood tests

parameter	Preoperative value	Postoperative value( 3 weeks)	Postoperative value(6weeks)
WBC	$2.99.\ 10^9 / L (4.0-10.010^9 / L)$	4.30. 10 <sup>9</sup> /L	$6.80.\ 10^9 / L$
Hemoglobin	8.2 g/L (115–150 g/L )	11.3 g/L	13.2 g/L
Platelets	80.000/mm <sup>3</sup> (110–320.)	130.000/mm <sup>3</sup>	240.000/mm <sup>3</sup>
Fibrinogen	1.83, g/L (2.00–4.00, g/L	2.13, g/L	3.22, g/L
INR	1.54 (0.85–1.50 )	1.35	1.10
Prothrombin time	18,2 sec (11–15).	15,1 sec	12,3 sec



FIG.1: CT scan images of a patient with Kasabach-Merritt syndrome associated with giant liver hemangioma

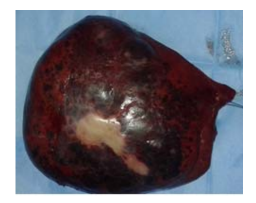


Fig.3: resected hemangioma

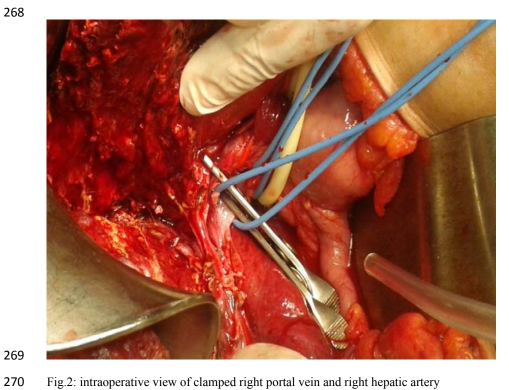


Fig.2: intraoperative view of clamped right portal vein and right hepatic artery