Combined therapy guided by multimodal imaging of fifteen retinal capillary hemangioblastomas in a monocular Von Hippel-Lindau syndrome case report

Ju Guo1†, Liping Du1†, Pengyi Zhou1, Xiaohong Guo2, Fangfang Dai2 and Xuemin Jin1*©

Abstract

Background: To report the multimodal imaging and treatment of fifteen retinal capillary hemangioblastomas (RCHs) associated with Von Hippel-Lindau syndrome in a monocular patient during a long-term following-up, which supply high-resolution exquisite SS-OCTA images (VG200; SVision Imaging, Ltd., Luoyang, China) and management experience about multiple RCHs.

Case presentation: A 34-year-old monocular male patient complained decreased visual acuity (20/100) without pain and redness in the left eye five years ago. Von Hippel-Lindau syndrome were diagnosed with genetic testing. He, his son and daughter all carried a heterozygosity missense variant c.499C > T (p. Arg167Trp) in the Hg19 gene, a VHL gene located in Chr3:10,191,506. Fifteen RCHs were identified by the application of multimodal imaging, which including fundus photo, fundus autofluorescence (FAF), B-scan ultrasonography (US), fluorescein fundus angiography (FFA), indocyanine green angiography (ICGA) and swept-source optical coherence tomography angiography (SS-OCTA). Transscleral cryotherapy and laser photocoagulation were performed to destroy the largest RCH with the size of 4 PD in diameter. Laser photocoagulation was conducted to seal the middle or tiny RCHs (< 1.5 PD) and their nourishing vessels. The retinal edema and exudative macular detachment were successfully relieved by intraocular injection of bevacizumab for 5 times. The RCHs in the left eye responded well to these treatments and best corrected visual acuity was 20/25 for three years. Three-month recall visits were recommended for him.

Conclusion: For multiple retinal capillary hemangioblastomas in monocular patients, precise combined therapy guided by multimodal imaging has a profound impact on the management of new and recurrent RCHs.

Keywords: Multiple retinal capillary hemangioblastomas, Swept-source optical coherence tomography angiography, Von Hippel-Lindau syndrome, Precise combined therapy, Case report

Background

VHL syndrome is an autosomal dominant inherited tumor syndrome caused by mutations of VHL gene, which is a tumour suppressor gene situated on chromosome 3p25 [1]. The mutations unbalance the expression of hypoxia-inducible factor, leading to tumours in the retina, central nervous system, kidneys and other organs [2]. Retinal capillary hemangioblastomas (RCHs) are benign tumours in nature and are always...
found in the periphery or juxtapapillary region. However, this syndrome usually leads to severe vision loss because of its serious complications, such as exudation, macular edema, retinal detachment, epiretinal membrane, and retinal and vitreous haemorrhage [3]. Therefore, it is necessary to timely detect and treat the RCHs in avoid of serious complications. In the past, the visualization of RCHs is mainly based on fluorescein fundus angiography (FFA), that can sketch the morphology of all VHL-related tumors in the retina. Swept-source optical coherence tomography angiography (SS-OCTA), a newly developed noninvasive ophthalmologic imaging technology, has been reported to provide a high-resolution representation for the true physical size of the RCHs [4, 5]. Nevertheless, the limited image range hamper the applications of SS-OCTA in clinical, so that it is important to get an accurate diagnosis by multimodal imaging of RCHs. Here, we report the multimodal imaging and treatment of fifteen retinal capillary hemangioblastomas (RCHs) associated with Von Hippel-Lindau syndrome in a monocular patient during the five-year following-up, which supply high-resolution exquisite SS-OCTA images and management experience about multiple RCHs. We obtained informed consent from the patient and his permission to report his case.

**Case presentation**

A 34-year-old male patient presented in our clinic on May 19, 2017, with a complaint of decreased visual acuity (20/100) in his left eye for several days. Laser photocoagulation had been performed at another hospital for the RCHs in his left eye half a year ago. He lost vision in his right eye after the surgical treatment of RCHs associated with VHL at the age of 30. The patient received multiple operations for cerebellar hemangioblastomas and adrenal pheochromocytoma when he was 25 years old. His father died from the complications of cerebellar hemangioblastomas associated with VHL. Genetic testing revealed that he, his son and daughter all carried a heterozygosity missense variant c.499C>T (p. Arg167Trp) in the Hg19 gene, a VHL gene located in Chr3:10,191,506 (Fig. 1).

The fundus photo, B-scan ultrasonography (US), fluorescein fundus angiography (FFA) and SS-OCT (single line scan mode) visualized eleven RCHs in the left eye (Fig. 2). Fundus photo showed the huge RCHs along with diffuse exudation around their nourishing vessels (No.1, 2, 7 RCHs in Fig. 2). FFA distinguished the middle and tiny ones in retina (No.3–6, 8-11RCHs in Fig. 2). B-scan ultrasonography provided the size, shape and location of the largest RCH (No.2 RCHs in Fig. 2). SS-OCT measured the size of the largest RCH and visualized its full-thickness structure (No.2 RCHs in Fig. 2).

---

**Fig. 1**  Pedigree chart and Sanger sequencing results of VHL gene variants using blood samples. The black square represents the patients who were clinically diagnosed with VHL. M/+ means heterozygous variants. The Sanger sequencing results show heterozygous missense variants in the VHL gene of patients II-1, III-1, and III-2
After twice consecutive monthly intravitreal ranibizumab (0.5 mg) injections and followed by prompt laser photocoagulation (within a-week day after every injection), obviously absorption of subretinal fluid (SRF) and visual acuity improvement (20/25) were observed (Fig. 3a-b). One year later, the patient complained vision loss again with exudative macular detachment and enlarged RCHs in nasal inferior retina due to unregular reexamination after combined therapy (approximately 3PD in diameter) (Fig. 4a). Four monthly ranibizumab intravitreal injections were performed, the reduced transient SRF and improved vision were noted (Fig. 3c-d), but the RCHs still enlarged rapidly (approximately 4PD in diameter) (Fig. 4b).

To control the enlarged RCHs and macular exudation, transscleral cryotherapy and intravitreal ranibizumab injection were performed under local anaesthesia in the left eye. One month after surgery, the largest RCH appeared to be pale and atrophied, accompanied by reduced exudation (Fig. 4c). Nine months after surgery, OCT images revealed that macular region remained a good structure and retinal exudation was reduced (Fig. 3e). Two years after surgery (November 6th, 2020), two newly developed RCHs temporal to macular were visualized by fundus examination using slit lamp due to the unregular reexamination. The multimodal imaging was recommended to this patient to find out all lesions in the retina in avoiding of serious complications. Fifteen RCHs were identified through detailed ophthalmological examination and multimodal imaging evaluation even if he had no complaint with vision loss (Fig. 5). The larger one with the size of 1.35*0.84*0.62 mm located in temporal to macular (No.5 RCH in Fig. 5) seemed to be exist in the first multimodal imaging evaluation, but it was so tiny that had been ignored during laser treatments. In addition, his irregular visit interrupted our observation and misled its diagnosis, so that No. 5 RCH grow rapidly during the two-year failure access. Then, three times continuous laser photocoagulations were performed around the RCHs and their nourishing vessels. Three months after laser therapy, the third multimodal image evaluation was conducted to observe the therapeutic effect and monitor active lesions (Fig. 6). The newly developed RCH (No.5 RCH in Fig. 5 and Fig. 6) responded well to laser treatments and best corrected visual acuity reaching 20/25. Following-up every three months were recommended for him to continuous monitor and treatment. The patient appreciated for the regained vision and maintained it for 3 years. Nevertheless, the financial burden of treatment may hinder his regular following-up.
Discussion and conclusions
Recent reports found missense mutations were the most common while protein-truncating mutations have a higher probability for a severe course [6, 7]. In this case, the patient carried heterozygosity missense variant c.499C > T (p. Arg167Trp) in the Hg19 gene, but his severe course of this disease hints us that every patient, regardless of the kind of mutation should be screened very thoroughly. The treatment of RCHs associated with Von-Hippel Lindau syndrome differs due to the size, location, complications of the hemangiomas. In the earlier stage of RCHs, careful observation is recommended for small RCHs (less than 0.5 mm in diameter) without the complications of diffuse exudation [8]. It is

Fig. 3 OCT (single line scan mode) image of exudation in the macular region during five-year following up. a SD-OCT showed macular detachment with exudative subretinal fluid (SRF) at his first visit on May 19th, 2017. b SD-OCT exhibited reduced SRF and exudation after twice anti-VEGF therapy and laser photocoagulation on September 10th, 2017. c SS-OCT revealed recurrent exudative macular detachment complicated with macular cystoid edema on August 17th, 2018, one year after the combined therapy. d SD-OCT visualized SRF and macular edema were completely absorbed with much exudation left after anti-VEGF therapy for four times on December 10th, 2018. e-f Nine months and two years after the cryotherapy (August 10th, 2019, November 6th, 2020), the macular region was still in good structure by the observations of SS-OCT Spectral-domain optical coherence tomography, SD-OCT Swept

source optical coherence tomography, SS-OCT subretinal fluid, SRF subretinal fluid

Fig. 4 Fundus photo of rapidly enlarged RCHs and the therapeutic effect of cryotherapy. a Fundus photo (taken in August 17th, 2018) showed an orange-red ellipse RCH with the diameter of approximately 3 PD was located in the nasal inferior retina, nourishing by two twisted and engorged vessels. b Fundus photo (taken in December 10th, 2018) showed the RCH located in the nasal inferior retina was rapidly enlarged to 4 PD in diameter and its nourishing vessels dilated with much exudation. c One month after cryotherapy, the RCH in the nasal inferior retina appeared to be pale and atrophied accompanied by reduced retinal exudation based on the fundus photo taken in January 28th, 2019. RCH retinal capillary hemangioblastomas, PD papillary diameter
usually effective for laser photocoagulation, photodynamic therapy (PDT), and external cryotherapy to treat small RCHs in peripheral retina (especially for RCH less than 1.5 mm in diameter) [8]. Brachytherapy or combined procedures have been recommended in the management of larger RCHs [9]. In cases with traction retinal detachment caused by RCHs, it is quite essential to treat by vitreoretinal surgery and tumour excision [10]. In addition, it has been demonstrated that the level of VEGF was elevated in the ocular fluids and in pathological specimens of eyes with VHL-associated RCH, so that the ophthalmologists have attempted to treat VHL-associated fundus lesions by the blockade of the VEGF signaling axis [11, 12]. Occasionally, intraocular anti-VEGF therapy has been reported to be useful in the reduction of exudation and lipid deposition without significant regression for VHL-related RCHs in recent case reports [13–15]. In this case, the effect of intravitreal anti-VEGF therapy on the exudation and lipid deposition was consistent with previous reports.

In this report, the largest RCH (No.2 RCH) was orange-red appearance with the size of 4 PD in diameter and accompany by diffuse exudation, which should be managed by invasive treatments, namely brachytherapy, PDT, or tumour resection. However, these treatments may increase the incidence of macular edema and vision loss [16]. Therefore, cryotherapy was conducted to destroy the largest RCH by the consideration of rehabilitating vision in this monocular patient. Laser photocoagulation was performed to seal the middle or tiny RCHs (< 1.5 PD) and their nourishing vessels. The retinal edema and exudative macular detachment were successfully relieved by intravitreal injection of bevacizumab for 5 times. Ultimately, the patient benefited a lot from the combined therapy and maintained a good vision for a long term.

SS-OCTA was a newly developed technology to visualize the microvasculature of the retina and choroid [17]. Previous research has demonstrated that SS-OCTA was advantage in detecting and measuring the size of RCHs in the posterior pole than other
ophthalmological technologies [18]. It has also been elected as most suitable imaging technology to monitor the RCHs due to it can provide a high-resolution image of hemangiomas and assessment for their size [5]. In this case, SS-OCTA was applied to identify tiny RCHs and figure out the structure and size of RCHs in posterior pole. What’s more, SS-OCTA depicted exquisite images of two RCHs in Fig. 5f. It contributed greatly in the long term following-up. FFA shows its advantage at sketching the contours of RCHs in peripheral retina, but can’t give a true assessment for their shape and texture features. Thus, it had been used to find out all potential RCHs in periphery in thrice multimodal image evaluations. What’s more, ICGA distinguished small RCHs in periphery from redundant fluorescent leakage points, due to its high protein binding rate. FAF can recognize the hypo-fluorescence middle sized RCHs from dilated vessels. B-scan ultrasonography give the information about the location and size of largest RCH, which rose above the retinal surface. In this case, all common ophthalmological imaging technologies were used to evaluate these fifteen RCHs, and their advantages were further discussed, which may have a profound impact on early detection of new and recurrent RCHs. However, the largest limitation of this case was the incident in the process of therapy due to his unregular reexamination.

In conclusion, it is safe and beneficial to utilize the combined therapy, namely transscleral cryotherapy, intravitreal ranibizumab injection and laser photocoagulation, guided by the multimodal imaging evaluation to treat multiple RCHs associated with Von-Hippel Lindau syndrome. In future following-up, it is important to apply the two magic weapons (multimodal imaging and combined therapy) to manage the RCHs.

**Abbreviations**

VHL: Von Hippel-Lindau; RCH: retinal capillary hemangioma; VEGF: vascular endothelial growth factor; FAF: Fundus autofluorescence; FFA: Fluorescein fundus angiography; ICGA: Indocyanine green angiography; SS-OCTA: Swept-source optical coherence tomography angiography.

**Acknowledgements**

Not applicable.
Authors' contributions
JG and PVZ analyzed and interpreted the patient data regarding the inspection results details and drafted the manuscript. XHG and FFD collected clinical data and helped draft the manuscript. LPD and XMJ participated in the analysis and interpretation of inspection results. All authors read and approved the final manuscript.

Funding
This work was supported by National Natural Science Foundation of China [81770914, 81970792, 82171040 and 81800832] and Medical Science and Technology Project of Health Commission of Henan Province [YXKC2020026]. These funding sources play important roles in study design, in the collection, analysis and interpretation of data, in the writing of the report, and in the decision to submit the paper for publication. National Natural Science Foundation of China,81970792,Liping Du ,82171040,Xuemin Jin ,Health Commision of Henan Province,YXKC2020026,Liping Du

Availability of data and materials
All data generated or analyzed during this study are included in this published article. The whole genome sequence result was provided in the supplementary file.

Declarations

Ethics approval and consent to participate
This case was approved by the Ethics Committee of the First Affiliated Hospital of Zhengzhou University (approved protocol number: 2021-KY-0736–002). Written informed consent for examinations and off-label treatments was obtained from the patient. In addition, written informed consent to participate in the genetic screening of his daughter and son was also obtained from the patient and his wife.

Consent for publication
Written informed consent was obtained from the patient for publication of this case report and any accompanying images. What’s more, written informed consent to publish the genetic testing result of his son and daughter was obtained from the patient and his wife. A copy of the written consent is available for review by the Editor of this journal.

Competing interests
The authors declare that they have no competing interests.

Author details
1Department of Ophthalmology, The First Affiliated Hospital of Zhengzhou University, Zhengzhou 450000, China. 2People’s Hospital of Zhengzhou University & Henan Eye Institute, Zhengzhou 450000, China.

Received: 22 August 2021 Accepted: 18 April 2022

Published online: 06 May 2022

References
1. Haddad NM, Cavallerano JD, Silva PS. Von hippel-lindau disease: a genetic and clinical review. Semin Ophthalmol. 2013;28(5–6):377–86.
2. Minervini G, Quaglia F, Tabaro F, Tosatto SCE. Insights into the molecular features of the von Hippel-Lindau-like protein. Amino Acids. 2019;51(10–12):1461–74.
3. Niemela M, Lemeta S, Sainio M, Rauma S, Pukkala E, Kere J, Bohling T, Laatikainen L, Jaaskelainen J, Summanen P. Hemangioblastomas of the retina: impact of von Hippel-Lindau disease. Invest Ophthalmol Vis Sci. 2000;41(7):1909–15.
4. Chun LY, Massamba N, Silas MR, Blair MP, Hariprasad SM, Skondra D. Use of optical coherence tomography angiography in the diagnosis of small retina lesions in Von Hippel-Lindau disease. Eye (Lond). 2020;34(12):2345–6.
5. Reich M, Glatz A, Boehringer D, Evers C, Daniel M, Buercher F, Ludwig F, Nuesse S, Lageze WA, Maloca PM, et al. Comparison of Current Optical Coherence Tomography Angiography Methods in Imaging Retinal Hemangioblastomas. Transl Vis Sci Technol. 2020;9(8):12.
6. Qiu J, Zhang K, Ma K, Zhou J, Gong Y, Cai L, Gong K. The Genotype-Phenotype Association of Von Hippel Lindau Disease Based on Mutation Locations: A Retrospective Study of 577 Cases in a Chinese Population. Frontiers in genetics. 2020;11:532588.
7. Reich M, Jaegle S, Neumann-Haefelin E, Klingler JH, Evers C, Daniel M, Buercher F, Ludwig F, Nuesse S, Kopp J, et al. Genotype-phenotype correlation in von Hippel-Lindau disease. Acta Ophthalmol. 2021;99(8):e1492–500.
8. Singh A, Shields C, Shields J. von Hippel-Lindau disease. Surv Ophthalmol. 2001;46(2):117–42.
9. Junker B, Schmidt D, Agostini HT. [Retinal angiomatosis. Ocular manifestation of von Hippel-Lindau disease]. Ophthalmologe. 2007;104(2):107–13.
10. Gaudric A, Krivosec V, Duguid G, Massin P, Giraud S, Richard S. Vitreoretinal surgery for severe retinal capillary hemangiomas in von hippel-lindau disease. Ophthalmology. 2011;118(1):142–9.
11. Chan CC, Vortmeyer AO, Chew EY, Green WR, Matteson DM, Shen DF, Linehan WM, Lubensky JA, Zhuang Z. VHL gene deletion and enhanced VEGF gene expression detected in the stromal cells of retinal angioma. Arch Ophthalmol. 1999;117(5):625–30.
12. Los M, Aarsman CJ, Terpstra L, Wittelsbeck-Post D, Lips CJ, Blijham GH, Voest EE. Elevated ocular levels of vascular endothelial growth factor in patients with von Hippel-Lindau disease. Ann Oncol. 1997;8(10):1015–22.
13. Agarwal A, Kumar N, Singh R. Intraretinal bevacizumab and feeder vessel laser treatment for a posteriorly located retinal capillary haemangioma. Int Ophthalmol. 2016;36(5):747–50.
14. Zubicco A, Andonegui J, Companes E, Plaza P, Tabuenca L, Mozio M. Combined treatment with intravitreal bevacizumab injection and photodynamic therapy for juxtapapillary retinal capillary haemangioma. J Fr Ophtalmol. 2020;43(4):e139–41.
15. von Buelow M, Pape S, Hoerauf H. Systemic bevacizumab treatment of a juxtapapillary retinal haemangioma. Acta Ophthalmol Scand. 2007;85(1):114–6.
16. Matsuou T, Himeki I, Ichimura K, Yanai H, Nose S, Mimura T, Miyoshi Y, Tsushima T. Long-term effect of external beam radiotherapy of optic disc hemangioma in a patient with von Hippel-Lindau disease. Acta Med Okayama. 2011;65(2):135–41.
17. Ferrara D, Waheed NK, Duker JS. Investigating the choriocapillaris and choroidal vasculature with new optical coherence tomography technologies. Prog Retin Eye Res. 2016;52:130–55.
18. Callaway NF, Muthyunjaya P. Widefield imaging of retinal and choroidal tumors. Int J Retina Vitreous. 2019;5(Suppl 1):49.