



ISSVA classification for vascular anomalies[©]

(Approved at the 20th ISSVA Workshop, Melbourne, April 2014)

Overview table

Vascular anomalies				
Vascular tumors	Vascular malformations			
	Simple	Combined °	of major named vessels	associated with other anomalies
<u>Benign</u>	<u>Capillary malformations</u>	<u>CVM, CLM</u>	<u>See details</u>	<u>See list</u>
<u>Locally aggressive or borderline</u>	<u>Lymphatic malformations</u>	<u>LVM, CLVM</u>		
<u>Malignant</u>	<u>Venous malformations</u> <u>Arteriovenous malformations*</u> <u>Arteriovenous fistula*</u>	<u>CAVM*</u> <u>CLAVM*</u> <u>others</u>		

° defined as two or more vascular malformations found in one lesion

* high-flow lesions

N.B. The classification tables do not list exhaustively all known vascular anomalies. Some rare "dermatologic" vascular anomalies will be found in dermatology textbooks.

The tumor or malformation nature or precise classification of some lesions is still unclear. These lesions appear in a separate provisional list.

[Abbreviations used](#)

For more details, click on the underlined links

Benign vascular tumors**Infantile hemangioma / Hemangioma of infancy**[see details](#)**Congenital hemangioma****Rapidly involuting (RICH) *****Non-involuting (NICH)****Partially involuting (PICH)****Tufted angioma * °****Spindle-cell hemangioma****Epithelioid hemangioma****Pyogenic granuloma (aka lobular capillary hemangioma)****Others****Locally aggressive or borderline vascular tumors****Kaposiform hemangioendothelioma * °****Retiform hemangioendothelioma****Papillary intralymphatic angioendothelioma (PILA), Dabska tumor****Composite hemangioendothelioma****Kaposi sarcoma****Others****Malignant vascular tumors****Angiosarcoma****Epithelioid hemangioendothelioma****Others**

* some lesions may be associated with thrombocytopenia and/or consumptive coagulopathy [see details](#)

° many experts believe that these are part of a spectrum rather than distinct entities

N.B. reactive proliferative vascular lesions are listed with benign tumors

Simple vascular malformations I

Capillary malformations (CM)

Cutaneous and/or mucosal CM (aka “port-wine” stain) [G](#)

CM with bone and/or soft tissue overgrowth

CM with CNS and/or ocular anomalies (Sturge-Weber syndrome)

CM of CM-AVM

[G](#)

CM of MICCAP (microcephaly-capillary malformation)

CM of MCAP (megalencephaly-capillary malformation-polymicrogyria)

Telangiectasia

Hereditary hemorrhagic telangiectasia (HHT) ([different types](#)) [G](#)

Others

Cutis marmorata telangiectatica congenita (CMTC)

Nevus simplex / Salmon patch / “angel kiss”, “stork bite”

Others

Simple vascular malformations II

Lymphatic malformations (LM)

Common (cystic) LM

Macrocystic LM

Microcystic LM

Mixed cystic LM

Generalized lymphatic anomaly (GLA)

LM in Gorham-Stout disease

Channel type LM

Primary lymphedema [\(different types\)](#)

G

Others

some lesions may be associated with thrombocytopenia and/or consumptive coagulopathy [see details](#)

clic on G to see genetics

Simple vascular malformations IIb

Primary lymphedema

Nonne-Milroy syndrome

[G](#)

Primary hereditary lymphedema

[G](#)

Lymphedema-distichiasis

[G](#)

Hypotrichosis-lymphedema-telangiectasia

[G](#)

Primary lymphedema with myelodysplasia

[G](#)

Primary generalized lymphatic anomaly

(Hennekam lymphangiectasia-lymphedema syndrome)

[G](#)

Microcephaly with or without chorioretinopathy,
lymphedema, or mental retardation syndrome

[G](#)

Lymphedema-choanal atresia

[G](#)

Simple vascular malformations III

Venous malformations (VM)

Common VM

[G](#)

Familial VM cutaneo-mucosal (VMCM)

[G](#)

Blue rubber bleb nevus (Bean) syndrome VM

Glomuvenous malformation (GVM)

[G](#)

Cerebral cavernous malformation (CCM) ([different types](#))

[G](#)

Others

some lesions may be associated with thrombocytopenia and/or consumptive coagulopathy [see details](#)

clic on [G](#) to see genetics

Simple vascular malformations IV

Arteriovenous malformations (AVM)

Sporadic

In HHT

[G](#)

In CM-AVM

[G](#)

Others

Arteriovenous fistula (AVF) (congenital)

Sporadic

In HHT

[G](#)

In CM-AVM

[G](#)

Others

Combined vascular malformations*		
CM + VM	capillary-venous malformation	CVM
CM + LM	capillary-lymphatic malformation	CLM
CM + AVM	capillary-arteriovenous malformation	CAVM
LM + VM	lymphatic-venous malformation	LVM
CM + LM + VM	capillary-lymphatic-venous malformation	CLVM
CM + LM + AVM	capillary-lymphatic-arteriovenous malformation	CLAVM
CM + VM + AVM	capillary-venous-arteriovenous malformation	CVAVM
CM + LM + VM + AVM	capillary-lymphatic-venous-arteriovenous m.	CLVAVM

* defined as two or more vascular malformations found in one lesion

Anomalies of major named vessels

(aka "channel type" or "truncal" vascular malformations)

Affect

- lymphatics
- veins
- arteries

Anomalies of

- origin
- course
- number
- length
- diameter (aplasia, hypoplasia, stenosis, ectasia / aneurysm)
- valves
- communication (AVF)
- persistence (of embryonal vessel)

Vascular malformations associated with other anomalies

Klippel-Trenaunay syndrome: CM + VM +/- LM + limb overgrowth

Parkes Weber syndrome: CM + AVF + limb overgrowth [G](#)

Servelle-Martorell syndrome: limb VM + bone undergrowth

Sturge-Weber syndrome: facial + leptomeningeal CM + eye anomalies
+/- bone and/or soft tissue overgrowth [G](#)

Limb CM + congenital non-progressive limb hypertrophy

Maffucci syndrome: VM +/- spindle-cell hemangioma + enchondroma

Macrocephaly - CM (M-CM / MCAP) [G](#)

Microcephaly - CM (MICCAP) [G](#)

CLOVES syndrome: LM + VM + CM +/- AVM + lipomatous overgrowth [G](#)

Proteus syndrome: CM, VM and/or LM + asymmetrical somatic overgrowth [G](#)

Bannayan-Riley-Ruvalcaba sd: AVM + VM +macrocephaly, lipomatous overgrowth [G](#)

Provisionally unclassified vascular anomalies

Verrucous hemangioma

Angiokeratoma

Multifocal lymphangioendotheliomatosis with thrombocytopenia / cutaneovisceral angiomatosis with thrombocytopenia (MLT/CAT)

Kaposiform lymphangiomatosis (KLA)

PTEN (type) hamartoma of soft tissue / "angiomatosis" of soft tissue

[G](#)

some lesions may be associated with thrombocytopenia and/or consumptive coagulopathy [see details](#)

clic on [G](#) to see genetics

Appendix 1 abbreviations used (excluding gene names)

AVF	arteriovenous fistula	HI	hemangioma of infancy / infantile hemangioma
AVM	arteriovenous malformation	IH	infantile hemangioma / hemangioma of infancy
CAT	cutaneovisceral angiomas with thrombocytopenia	INR	international normalized ratio
CAVM	capillary arteriovenous malformation	JPHT	juvenile polyposis hemorrhagic telangiectasia
CCM	cerebral cavernous malformation	KHE	kaposiform hemangioendothelioma
CLAVM	capillary lymphatic arteriovenous malformation	KLA	kaposiform lymphangiomatosis
CLOVES	congenital lipomatous overgrowth, vascular malformations, epidermal nevi, skeletal/scoliosis and spinal abnormalities	KMP	Kasabach-Merritt phenomenon,
CLM	capillary lymphatic malformation	LM	lymphatic malformation
CLVAVM	capillary lymphatic venous arteriovenous malformation	LVM	lymphatic venous malformation
CLVM	capillary lymphatic venous malformation	MCAP	megalencephaly-capillary malformation-polymicrogyria
CM	capillary malformation	M-CM	macrocephaly-capillary malformation
CM-AVM	capillary malformation-arteriovenous malformation	MICCAP	microcephaly-capillary malformation
CMTC	cutis marmorata telangiectatica congenita	MLT	Multifocal lymphangioendotheliomatosis with thrombocytopenia
CNS	central nervous system	NICH	non-involuting congenital hemangioma
CVAVM	capillary venous arteriovenous malformation	PHACE	posterior fossa malformations, hemangioma, arterial anomalies, cardiovascular anomalies, eye anomalies
CVM	capillary venous malformation	PILA	papillary intralymphatic angioendothelioma
DIC	disseminated intravascular coagulopathy	PICH	partially involuting congenital hemangioma
GLA	generalized lymphatic anomaly	RICH	rapidly involuting congenital hemangioma
GSD	Gorham-Stout disease	TA	tufted angioma
GVM	glomuvenous malformation	VM	venous malformation
HHT	hereditary hemorrhagic telangiectasia	VMCM	venous malformation cutaneo mucosal

Appendix 2-a

causal genes of vascular anomalies

Capillary malformations (CM)	
Cutaneous and/or mucosal CM (aka “port-wine” stain)	GNAQ
CM with bone and/or soft tissue hyperplasia	
CM with CNS and/or ocular anomalies (Sturge-Weber syndrome)	GNAQ
CM of CM-AVM	RASA1
Telangiectasia	
Hereditary hemorrhagic telangiectasia (HHT)	
<i>HHT1</i>	ENG
<i>HHT2</i>	ACVRL1
<i>HHT3</i>	
<i>JPHT</i> (juvenile polyposis hemorrhagic telangiectasia)	SMAD4
Others	
Cutis marmorata telangiectatica congenita (CMTC)	
Nevus simplex / Salmon patch	
Others	

Appendix 2-b

causal genes of vascular anomalies

Lymphatic malformations (LM)

Primary lymphedema

Nonne-Milroy syndrome	FLT4 / VEGFR3
Primary hereditary lymphedema	VEGFC
Primary hereditary lymphedema Connexin 47	GJC2 /
Lymphedema-distichiasis	FOXC2
Hypotrichosis-lymphedema-telangiectasia	SOX18
Primary lymphedema with myelodysplasia	GATA2
Primary generalized lymphatic anomaly (Hennekam lymphangiectasia-lymphedema syndrome)	CCBE1
Microcephaly with or without chorioretinopathy, lymphedema, or mental retardation syndrome	KIF11
Lymphedema-choanal atresia	PTPN14

Appendix 2-c

causal genes of vascular anomalies

Venous malformations (VM)	
Common VM	TIE2 somatic
Familial VM cutaneo-mucosal (VMCM)	TIE2
Blue rubber bleb nevus (Bean) syndrome VM	
Glomuvenous malformation (VM with glomus cells)	Glomulin
Cerebral cavernous malformation (CCM)	
CCM1	KRIT1
CCM2	Malcavernin
CCM3	PDCD10

Appendix 2-d

causal genes of vascular anomalies

Arteriovenous malformations (AVM)	
Sporadic	
In HHT	
<i>HHT1</i>	ENG
<i>HHT2</i>	ACVRL1
<i>JPHT (juvenile polyposis hem. telangiect.)</i>	SMAD4
In CM-AVM	RASA1
Arteriovenous fistulas (AVF)	
Sporadic	
In HHT	
<i>HHT1</i>	ENG
<i>HHT2</i>	ACVRL1
<i>JPHT (juvenile polyposis hemorrhagic telangiectasia)</i>	SMAD4
In CM-AVM	RASA1

Appendix 2-e

causal genes of vascular anomalies

Vascular malformations associated with other anomalies	
Klippel-Trenaunay syndrome	
Parkes Weber syndrome	RASA1
Servelle-Martorell syndrome	
Sturge-Weber syndrome	GNAQ
Limb CM + congenital non-progressive limb overgrowth	
Maffucci syndrome	
Macrocephaly - CM (M-CM or MCAP)	PIK3CA
Microcephaly - CM (MICCAP)	STAMB P
CLOVES syndrome	PIK3CA
Proteus syndrome	AKT1
Bannayan-Riley-Ruvalcaba syndrome	PTEN

Appendix 2 -f

causal genes of vascular anomalies

Provisionally unclassified vascular anomalies

Verrucous hemangioma

Multifocal lymphangioendotheliomatosis with thrombocytopenia / cutaneovisceral angiomatosis with thrombocytopenia (MLT/CAT)

Kaposiform lymphangiomatosis (KLA)

PTEN (type) hamartoma of soft tissue / "angiomatosis" of soft tissue **PTEN**

some lesions may be associated with thrombocytopenia and/or consumptive coagulopathy [see details](#)

Appendix 3

infantile hemangioma

Pattern	Different types
<ul style="list-style-type: none">- focal- multifocal- segmental- indeterminate	<ul style="list-style-type: none">- superficial- deep- mixed (superficial + deep)- reticular / abortive / minimal growth- others

Association with other lesions	
PHACE association / syndrome	Posterior fossa malformations, Hemangioma, Arterial anomalies, Cardiovascular anomalies, Eye anomalies, sternal clefting and/or supraumbilical raphe
LUMBAR (SACRAL, PELVIS) association / syndrome	Lower body hemangioma, Urogenital anomalies, Ulceration, Myelopathy, Bony deformities, Anorectal malformations, Arterial anomalies, and Renal anomalies

possibly associated with platelet count / coagulation disorders

Anomalies	Hematological disorders
Tufted angioma Kaposiform hemangioendothelioma	Profound and sustained thrombocytopenia with profound hypofibrinogenemia, consumptive coagulopathy and elevated D-dimer (Kasabach-Merritt phenomenon)
Rapidly involuting congenital hemangioma	Transient mild/moderate thrombocytopenia, +/- consumptive coagulopathy and elevated D-dimer
Venous malformations / Lymphatic-venous malformations	Chronic localized intravascular coagulopathy with elevated D-dimer, +/- hypofibrinogenemia, and +/- moderate thrombocytopenia (may progress to DIC after trauma or operation)
Lymphatic malformations	Chronic localized intravascular coagulopathy with elevated D-dimer and +/- mild to moderate thrombocytopenia <i>(consider Kaposiform lymphangiomatosis)</i> (may progress to DIC after trauma or operation)
Multifocal lymphangioendotheliomatosis with thrombocytopenia / Cutaneovisceral angiomatosis with thrombocytopenia	Sustained, fluctuating, moderate to profound thrombocytopenia with gastrointestinal tract bleeding or pulmonary hemorrhage
Kaposiform lymphangiomatosis	Mild to Moderate thrombocytopenia, +/- hypofibrinogenemia, and D-dimer elevation