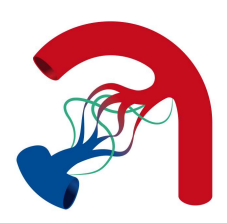


for rare or low prevalence complex diseases

: Network

Vascular Diseases (VASCERN)



Lymphatic malformation

Final Approved Patient Pathway by the Vascular Anomalies (VASCA) Working Group - 29/04/2020

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Disclaimer

- This document is an opinion statement reflecting strategies put forward by experts and patient representatives involved in the Vascular Anomalies (VASCA) Rare Disease Working Group of VASCERN.
- It is preferable that patients be evaluated in a multidisciplinary center specialized in the diagnosis and management of vascular anomalies.
- This pathway is issued on 29/04/2020 and will be further validated and adjusted as needed.
- Multidisciplinary team should re-evaluate treatment decisions regularly
- Responsibility for care of individual patients remains with the treating physician.

Dieses Dokument wurde am 05.12.2022 von Dr. Friedrich Kapp für das Zentrum für Gefäßfehlbildungen (Uniklinik Freiburg) für das FZSE Freiburg geprüft und kommentitiert. Eine gemeinsame Überprüfung und Diskussion wird unabhängig hiervon in VASCA-WG erfolgen.

Dieses Dokument wurde am 15.11.2023 von Dr. Friedrich Kapp für das Zentrum für Gefäßfhelbildungen (Uniklinik Freiburg) für das FZSE Freiburg geprüft und kommentiert. Eine gemeinsame Überprüfung und Diskussion wir unabhängig hiervon in der VASCA-WG erfolgen.

Dieses Dokument wurde am 10.10.2024 von Dr. Friedrich Kapp für das Zentrum für Gefäßfhelbildungen (Uniklinik Freiburg) für das FZSE Freiburg geprüft und kommentiert. Eine gemeinsame Überprüfung und Diskussion wir unabhängig hiervon in der VASCA-WG erfolgen.





Gathering the best expertise in Europe to provide accessible cross-border healthcare to patients with rare vascular diseases



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Lymphatic malformation Diagnostic Work-Up: Overview

Suspected lymphatic malformation

Possible clinical signs or symptoms:

- Prenatal detection of a cyst
- Onset most often in early childhood
- Gradual swelling of a (subcutaneous) non-solid, non-compressible mass
 - · Bluish hue on the skin
 - Vesicles
 - Recurrent lymph leak/oozing/ bleeding
 - Frequent skin infections and inflammations in the mass
- Protein loosing enteropathy / pleural- pericardial or abdominal fluid effusion or cysts
 - Most often no pain
 - Incidental finding
 - Edema
 - Doppler ultrasound to confirm diagnosis & exclude flow in the lesions
 - MRI if diagnosis unclear or prior to surgery or sclerotherapy
 - CT if suspicion of bone involvement (Gorham-Stout disease)
 - Lymphatic scintigraphy if suspicion of lymphedema
 - Mucosal investigation with endoscopy
- Lymphangiography if suspicion of channel type LM or pleural-, pericardial- or peritoneal chylus effusion
- · Coagulation work-up (Chronic localized intravascular coagulopathy?) Kasabach-Merritt Phenomenon (KMP)
 - Biopsy or FNAC occasionally needed for differential diagnosis *

*Histology is helpful for differential diagnosis: LMs are composed of thinwalled, dilated irregular channels which often appear empty or contain pale eosinophilic amorphous material. Superficial LM may be associated with overlying epidermal hyperplasia and hyperkeratosis. Immunostains for lymphatic markers are helpful in differentiating LM from other malformations. PROX1. VEGFR-3. D2-40 (podoplanin) and LYVE-1 all label lymphatic endothelium. * Cytology and cyst chemistry is used to rule out cysts of other origin such as pseudocysts from parenchymal organs

Somatic PIK3CA mutation is associated to most LM Simple/Common LM LM part of LM in Gorham-Stout Channel type LM **Syndromic** Primary lymphedema Generalized lymphatic disease (GSD) (i.e. PROS) Kommentar FK 11/2023 anomaly (GLA) and akutelle Bezeichnung ist central conducting lymphatic anomaly **Kaposiform** (CCLA) Lymphangiomatosis Please see pathway for *Syndromic* Please see pathway from (KLA) vascular malformations **VASCERN PPL-WG** Clinical Associated LEGEND: **Investigations** Treatment Particular cases

evaluation

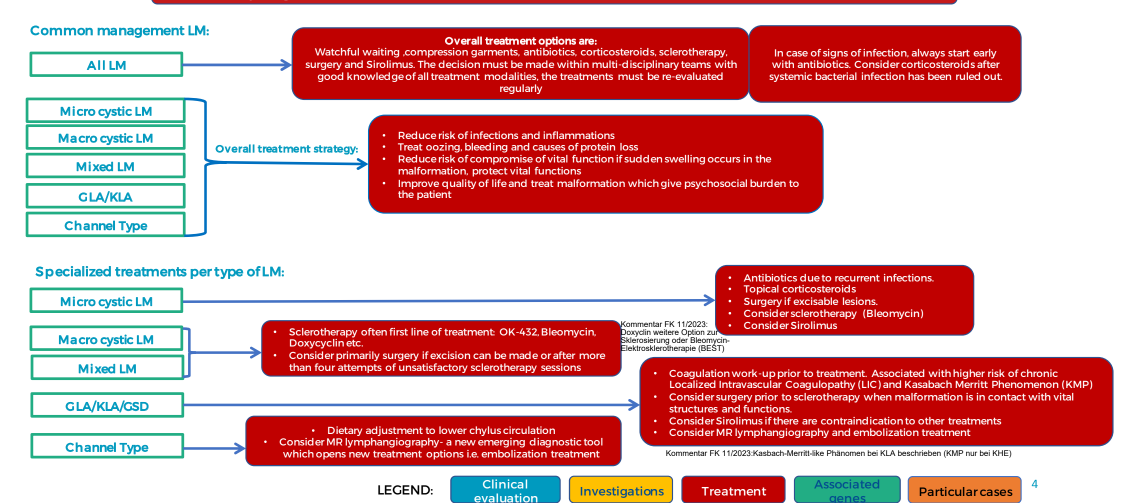






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Lymphatic malformation: Management/Treatment Overview

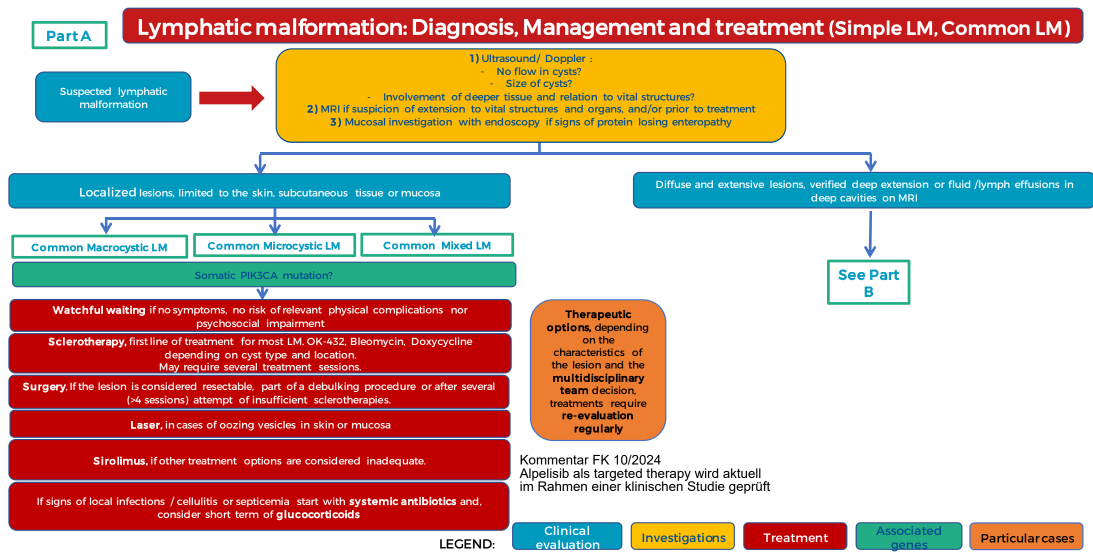








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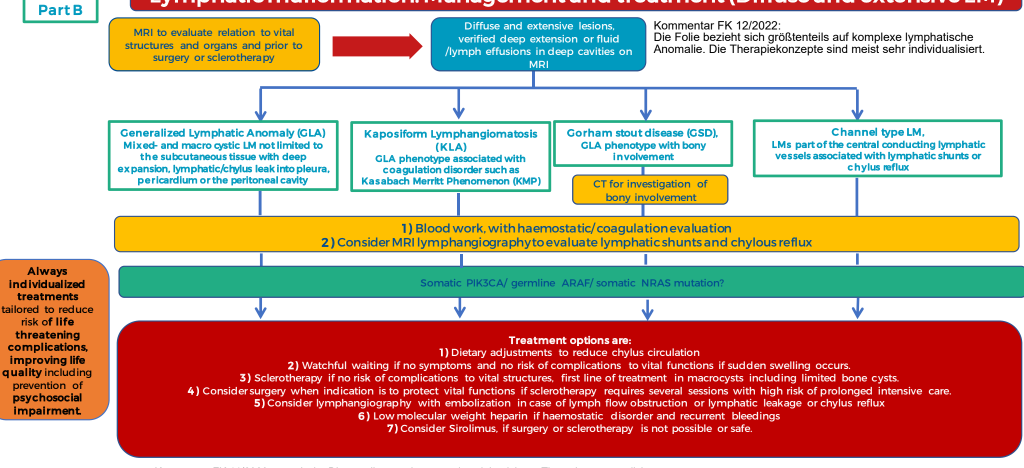


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Lymphatic malformation: Management and treatment (Diffuse and extensive LM)



Kommentar FK 11/2023: genetische Diagnostik anstreben, um eine zielgerichtete Therapie zu ermöglich

LEGEND:

Clinical evaluation

Investigations

Treatment

Associated genes

Particular cases



VASCERN

Gathering the best expertise in Europe to provide accessible cross-border healthcare to patients with rare vascular diseases



VASCERN, the European Reference Network on Rare Multisystemic Vascular Diseases, is dedicated to gathering the best expertise in Europe in order to provide accessible cross-border healthcare to patients with rare vascular diseases (an estimated 1.3 million concerned). These include arterial diseases (affecting aorta to small arteries), arterio-venous anomalies, vascular malformations, and lymphatic diseases.

VASCERN currently consists of 30 highly specialised multidisciplinary Healthcare Providers (HCPs) from 11 EU Member States and of various European Patient Organisations and is coordinated in Paris, France.

Through our 5 Rare Disease Working Groups (RDWGs) as well as several thematic WGs and the ePAG - European Patient Advocacy Group, we aim to improve care, promote best practices and guidelines, reinforce research, empower patients, provide training for healthcare professionals and realise the full potential of European cooperation for specialised healthcare by exploiting the latest innovations in medical science and health technologies.

More information available at: https://vascern.eu

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