What Does This Mean for Patients?

Reliable Information

Access to clear and up-to-date information about LMS and the latest research developments.

Community and Connection

Opportunities to connect with other LMS patients and families through support groups and online forums, fostering a sense of belonging and shared experience.

Knowledge

Resources to help you advocate for yourself and make informed decisions about your treatment and care.

Hope for the Future

By funding research and fostering collaboration, we are actively working towards better treatments and improved outcomes for LMS patients.



Contact Us

https://lmsruk.or: 67-169 Great Portland Street 5th Floor, London. W1W SPF

Who are LMSR UK?

We are a newly established organisation founded by LMS patients and their loved ones. Driven by the urgent need for better treatments and outcomes, we are dedicated to making a real difference in the lives of those affected by LMS.

Our mission is to support research, provide education, and foster a strong community for LMS patients and their families across the UK.

RESEARCH UK

Together, we will find a cure.



LMS Research UK

Registered UK Charity 1210455

How To Get Involved?

Join Our Healthcare Professional Network

Work together with us to strengthen a UK network of healthcare professionals interested in LMS linked Soft Tissue Sarcoma research.

Join Our Community

Connect with us through our support groups, online forums, and social media channels.

Raise Awareness

Share Information about LMS and LMSR UK with your network, helping to increase understanding and support.

Fundraise or Donate

Donate to our JustGiving page or organise fundraising events to support us.

Volunteer

Offer your time and skills to help us achieve our mission.

Our Aims

Establish a UK Network of LMS Experts

We aim to create a network of researchers and experts to coordinate and focus research efforts, encourage collaboration, and share best practices, ultimately leading to improved outcomes for LMS patients.

Educate and Inform

We provide up-to-date, accessible information about LMS through leaflets, our 'website, social media, and patient information sessions.

Provide Support

We offer a safe and supportive community for those affected by LMS through local and online support groups and forums.

Raise Awareness

We actively work to increase awareness of LMS within the community and among medical professionals, advocating for better diagnosis and treatment options.

Sponsor Research

We are dedicated to funding research into the prevention, diagnosis, and treatment of LMS, with the goal of improving patient outcomes and developing the next generation of medical experts and researchers.

What we are working on:

- Setting up a national (virtual) tissue bank for all LMS Tissue which can be accessible for all UK LMS researchers.
- **2.** Landscape mapping of the UK Uterine Sarcomas practice (to understand the differences in regional approaches)
- **3.** A prospective Radiomics trial using MRIs and Circulating Tumour DNA to support early diagnosis.
- **4.** Support towards an Adjuvant Chemotherapy Trial
- **5.** Pushing for Trabectadin and Doxorubicin to be the first line standard of care for the UK.
- **6.** Pushing forward on the UK Government Rare Cancer Bill to make LMS more prominent with a positive impact on UK LMS patients.

For More Info, Visit:



Or Email: admin@lmsruk.org if you are interested to work together.