WORLD LYMPHEDEMA DAY 2020

An overview with suggestions on how to celebrate and raise awareness on March 6th and beyond

A collaboration among patient representatives

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The Netherlands

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Sweden
Associations

Denmark
- Dansk Lymfødem Forening

Italy
- Lymphido

The Netherlands
- Nederlands Netwerk voor Lymfoedeemen & Lipoedeem

Portugal
- Associação Nacional de Doentes Linfáticos

Spain
- Asociación Galega de Linfedema, Lipedema y Patologías Vasculares Periféricas

Sweden
- Svenska Ödemförbundet
OUR GOALS for WORLD LYMPHEDEMA DAY

Put lymphedema on the radar of politicians
- make sure politicians realise the scope of the problem, how many patients are affected, know that lymphedema is a chronic condition, the issues surrounding getting a diagnosis and adequate treatment

Increase membership
- make sure patients know about their national patient association and that they understand the power in numbers, the support we can offer and work we are doing to create changes

Engage clinicians
- make sure they know what lymphedema is, signs and symptoms of lymphedema, difference between primary and secondary lymphedema, the risk of secondary lymphedema after cancer treatment, direct patients to patient associations and support groups

Raise awareness with the public
- about lymphedema, what is it, how to identify it, what to do if you think you have it, direct to more information. What is World Lymphedema Day?
Let’s get to work

Collation of information
Gather information from patient associations, other resources

Sharing ideas
Discussing with other patient associations and patient advocates

Disseminating overview
Share with wider community, get feedback on uptake

Creating overview
Overview to be written and checked by parties involved

Coordinate
Coordinate overlapping activities across countries, PAs, advocates

Monitor
Start monitoring activities and collate data

March 6th

ROADMAP to WORLD LYMPHEDEMA DAY

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ACTIVITIES

Online activities include use of social media and landing page. Use assets and locally generated content for shareability.

Events organized by the patient associations such as seminars, workshops, "drop-ins", a table in a hospital/clinic, in the street or at other venues relevant to get into contact with the public or with clinicians, flash mobs, catwalks.

“Events” organized by the patients such as a walk, lunch, a fundraiser, patient-to-patient meeting or encounter. Using social media to share content or personal testimonial. Start a blog, an Instagram account, public Facebook profile or Twitter.

Events organized/supported by clinicians, health care providers and other organisations.
**TARGET areas**

**Social Media**
Use the outlets of the patient associations to spread awareness. Encourage patients to share content from these to ensure data collection and link back to PA.

**Website/e-mails**
Landing page on PA website to link to in emails – ensure links to other resources national/international. Use PA emailing list to reach members not on social media.

**Patients**
Patient collaboration – especially patient advocates with big following, board members, those with access to the press. Get them to tell their story. Organize local events.

**Press**
Use PR and contacts to reach the wider public. Circulate stories for higher reach.
USE of POPULAR SOCIAL MEDIA

**Facebook**
Most patient associations have a Facebook public page and associated open, closed or private groups. Best for reaching the patient “masses” and their friends.

**Instagram**
Some patient associations have a profile. Most patient advocates use it. Best for reaching younger patients, parents with children with LE, patient advocates, “hands-on” clinicians and companies.

**Twitter**
Some patient associations have a Twitter account. Best for reaching clinicians, researchers, hospitals, politicians.

**YouTube**
Best for storing video material to be linked to in other platforms.

**Pinterest**
Unexplored. Potentially to make use of SEO on keywords such as lymphedema, swollen legs, puffy feet, edema. Potential to reach undiagnosed patients.

**LinkedIn**
Unexplored. Potentially used for reaching representatives of companies and clinicians.

**Communities**
Forums and communities like Rareconnect.
• Create landing page on local patient association website to list ideas for patients and share what patient association is doing to celebrate WLD in local language

• Create event on Facebook page

• Send email to members and direct them to the page, ask them to follow patient association(s) on social media platforms and share the content on public or in private messages to contacts

• Send mail/e-mail to politicians, communities, boards

• Throughout the day or week on social media
  • Post top facts about lymphedema
  • Post testimonials of patients
  • Post information about the difficulty in getting the correct treatment
  • Make sure to tag and use appropriate hashtags

• Link to WHO petition
ACTIVITIES

• Organise a run/walk
• Set up in local hospital or on streets
  • Table, flyers, registration forms, banner, volunteers to talk
• Support the manifesto (local language), share it and read it in public areas
  • Send to/invite press, engage a spokesperson or public figure
• Flash mobs or catwalks in public places
• Support publication of articles or dissemination of surveys for research
• Submit abstracts to conferences
• Seminars
  • Inviting patients, clinicians, politicians, public service workers
• Workshops/hands-on/coffee meeting for patients
• Send mail to politicians, communities, boards
• Ask patients to sign WHO petition
• Log your event on the website
  • https://lymphaticnetwork.org/wld/create/

EVENTS - PA

[Images of events: Reading the manifesto in Spain, Out on the streets with banners and flyers in Norway, The Netherlands: Photo shot with patients with lymphedema, Italy: Left and right patients with secondary and primary lymphedema. Centre patient with congenital melanocytic nevus]

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ACTIVITIES

• Organise a run/walk/coffee morning/lunch among friends and family.
• Bring cake to work and ask co-workers to read a flyer to get one
• Consider raising funds for patient association or ask friends and family to become members to show support
• Start a blog or Instagram account about their life as a patient with lymphedema – follow others!
• Share their story on other outlets like LE&RN, in forums like Rareconnect or via the patient associations website or social media accounts
  
  • https://lymphaticnetwork.org/living-with-lymphedema/your-stories/
  • https://www.rareconnect.org

• Attend events – get out and meet other patients
• Visit local hospitals and GP practices and hand out flyers
• Volunteer skills and time to the patient association
• Wear blue, a teal/blue ribbon, a specially made t-shirt
• Create business cards with your instahandle or other social media profile
• #flauntyourcompression – show the world your compression and remember to use hashtags
• Decorate your profile pictures with a twibbon. https://twibbon.com/Support/world-lymphedema-day
• Sign the petition
  
SOCIAL MEDIA
What can the patients do online?

Facebook personal profile
- Add profile frame
- Change profile picture/cover picture
- Write a personal testimonial
- Share content from patient associations

Patients groups and forums
- Patients should be encouraged to share their tips on how to manage their lymphedema, living with lymphedema
- Answer questions, encourage other patients
- Invite other patients to the patient groups
- Join Rareconnect (for primary patients)

Twitter & YouTube
- Write post
- Share content

Instagram
- If patient has a profile dedicated to lymphedema then share testimonial, content and other patient stories
- If patient has non-lymphedema related profile consider to add a post on March 6
- A patient can consider to create their own Instagram profile sharing their life with lymphedema

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HASHTAGS

International
#worldlymphedemaday, #wld2020, #lymphedema #lymphoedema #lymphedemaawereness
#nevergiveup #flauntyourcompression #rockthesock #primarylymphedema #secondarylymphedema

National
Local language words for lymphedema, local language words for lymphedema day or world lymphedema day etc. Check other patients use of commonly used hashtags in your country

Tagging
Don’t forget to tag associations, organisations, clinicians and companies, tag each other

Share
Make sure content can be shared in public
Clinicians and Researchers

How to engage them?

Patients

• Ask to leave flyers/leaflets/factsheet in the GP practice
• Make sure you discuss your lymphedema at the visit...even just to say all is well – make sure they remember how it looks!
• Take the time to speak to the receptionist and nurse practitioner – use a leaflet to start the conversation and explain why you have lymphedema. Tell them to refer patients to the patient associations or Facebook patient groups – help a fellow patient

Patient associations

• Email contacts about your activities for WLD and ask for their support in sharing and celebrating the day
• Tag organisations/societies/key opinion leaders on posts on social media
• Make sure they know you as a patient association so they can refer patients to you
• Support research and dissemination of surveys

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Many excellent videos are already available on YouTube or websites.

Factsheets are available and could be translated into local language for easier usage.

The national and international websites are packed with information which can be shared and used.

We lack press release material to copy and translate in local language.

Many excellent videos are already available on YouTube or websites.
### ASSETS - WEBSITES

**English language**

<table>
<thead>
<tr>
<th>Organisation/Association (English)</th>
<th>Web address</th>
</tr>
</thead>
<tbody>
<tr>
<td>LE&amp;RN</td>
<td><a href="https://lymphaticnetwork.org/">https://lymphaticnetwork.org/</a></td>
</tr>
<tr>
<td>LE&amp;RN WLD</td>
<td><a href="https://lymphaticnetwork.org/wld/get-involved/">https://lymphaticnetwork.org/wld/get-involved/</a></td>
</tr>
<tr>
<td>Canadian Lymphedema Framework</td>
<td><a href="https://canadalymp.ca/">https://canadalymp.ca/</a></td>
</tr>
<tr>
<td>International Lymphoedema Framework</td>
<td><a href="https://www.lympho.org/">https://www.lympho.org/</a></td>
</tr>
<tr>
<td>Lymphoedema Support Network</td>
<td><a href="https://www.lymphoedema.org/">https://www.lymphoedema.org/</a></td>
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<tr>
<td>Lymphoedema Ireland</td>
<td><a href="https://www.lymphireland.com/">https://www.lymphireland.com/</a></td>
</tr>
<tr>
<td>L-W-O</td>
<td><a href="https://lymph-what-oedema.com/">https://lymph-what-oedema.com/</a></td>
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</tbody>
</table>
# ASSETS - WEBSITES

**Local language**

<table>
<thead>
<tr>
<th>Patient Association</th>
<th>Language</th>
<th>Web address</th>
</tr>
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<tbody>
<tr>
<td>andLINFA</td>
<td>Portuguese</td>
<td><a href="http://www.andlinfa.pt">www.andlinfa.pt</a></td>
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<tr>
<td>AGL</td>
<td>Spanish</td>
<td><a href="http://www.linfedemagalia.com">www.linfedemagalia.com</a></td>
</tr>
<tr>
<td>AVML</td>
<td>French</td>
<td><a href="http://www.avml.fr">www.avml.fr</a></td>
</tr>
<tr>
<td>DALYFO</td>
<td>Danish</td>
<td><a href="http://www.dalyfo.dk">www.dalyfo.dk</a></td>
</tr>
<tr>
<td>KIF11</td>
<td>German</td>
<td><a href="http://www.kif11kids.com">www.kif11kids.com</a></td>
</tr>
<tr>
<td>Lymphido</td>
<td>Italian</td>
<td><a href="http://www.lymphido.it">www.lymphido.it</a></td>
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<tr>
<td>NLNet</td>
<td>Dutch</td>
<td><a href="http://www.nlnet.nl">www.nlnet.nl</a></td>
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<tr>
<td>NLF</td>
<td>Norwegian</td>
<td><a href="http://www.nllf.no">www.nllf.no</a></td>
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<tr>
<td>SOF</td>
<td>Swedish</td>
<td><a href="http://www.svenskaodemforbundet.se">www.svenskaodemforbundet.se</a></td>
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</tbody>
</table>

**Your own local language resources!**

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<tr>
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## ASSETS- VISUAL CONTENT

<table>
<thead>
<tr>
<th>Video</th>
<th>Link</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lymphedema from a child’s perspective</td>
<td><a href="https://youtu.be/CrzyAaj655U">https://youtu.be/CrzyAaj655U</a></td>
</tr>
<tr>
<td>An Overview of Pediatric and Primary Lymphedema</td>
<td><a href="https://youtu.be/zcYSmxX_RPC">https://youtu.be/zcYSmxX_RPC</a></td>
</tr>
<tr>
<td>My lymphedema in a few words</td>
<td><a href="https://youtu.be/lf75ueX-2aI">https://youtu.be/lf75ueX-2aI</a></td>
</tr>
<tr>
<td>The lymphatic system and lymphedema: how it works</td>
<td><a href="https://youtu.be/HEiW0bE20wI">https://youtu.be/HEiW0bE20wI</a></td>
</tr>
<tr>
<td>LE&amp;RN WLD 2020</td>
<td><a href="https://youtu.be/PMZw0N9p2Nk">https://youtu.be/PMZw0N9p2Nk</a></td>
</tr>
<tr>
<td>You are never alone – Lymphedema and Lipedema Community message</td>
<td><a href="https://youtu.be/zRWfViRp6dY">https://youtu.be/zRWfViRp6dY</a></td>
</tr>
<tr>
<td>What is lymphoedema (with Denise Hardy)</td>
<td><a href="https://youtu.be/6nzwPEqLn8E">https://youtu.be/6nzwPEqLn8E</a></td>
</tr>
<tr>
<td>A collection of patient stories with challenges and top tips</td>
<td><a href="https://www.youtube.com/playlist?list=PLekVty32TyUm5z2d2rW4WejrgbdB7Rk6w">https://www.youtube.com/playlist?list=PLekVty32TyUm5z2d2rW4WejrgbdB7Rk6w</a></td>
</tr>
<tr>
<td>What is Lymphoedema</td>
<td><a href="https://vimeopro.com/ehdm/lsn-18/video/299859108">https://vimeopro.com/ehdm/lsn-18/video/299859108</a></td>
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### Your own local language resources!

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</table>
There is not much easy to use or copy press release material available

- An example from Sweden

- LE&RN website

Please share what you do with other patient associations
# ASSETS - FACT SHEETS

<table>
<thead>
<tr>
<th>Fact Sheet</th>
<th>Website</th>
</tr>
</thead>
<tbody>
<tr>
<td>Do's and Don'ts Fact Sheet for clinicians</td>
<td><a href="http://www.vascern.eu">www.vascern.eu</a></td>
</tr>
<tr>
<td>Teenage Lymphoedema, Early Childhood Lymphoedema</td>
<td><a href="http://www.lymphoedema.org">www.lymphoedema.org</a></td>
</tr>
<tr>
<td>10 Things….a range of Fact Sheets</td>
<td><a href="https://lymphaticnetwork.org/resource-downloads">https://lymphaticnetwork.org/resource-downloads</a></td>
</tr>
</tbody>
</table>
DATA which can be COLLECTED

GOALS
#patients, clinicians, public reached
#of new members in your association

AWARENESS
#PR uptake, press attendance at event

EMAILS/WLD LANDING PAGE
#emails send to members, clinicians, public sector, politicians.
#visits to landing page

COST
Cost for events, gadgets, flyers, etc.

ACTIVITIES
#volunteers engaged in supporting the work of the patient associations.
#activities done, #attendance

SOCIAL MEDIA
#posts/content shared, likes, shares, comments, visits, click-throughs, new followers, downloads, tags, testimonials

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THANK YOU

We hope you liked our suggestions and ideas. Please share with us what you do so we can exchange best practice across Europe.

We wish you all the best for a successful World Lymphedema Day!