

Building Community • Giving Voice • Advancing Care

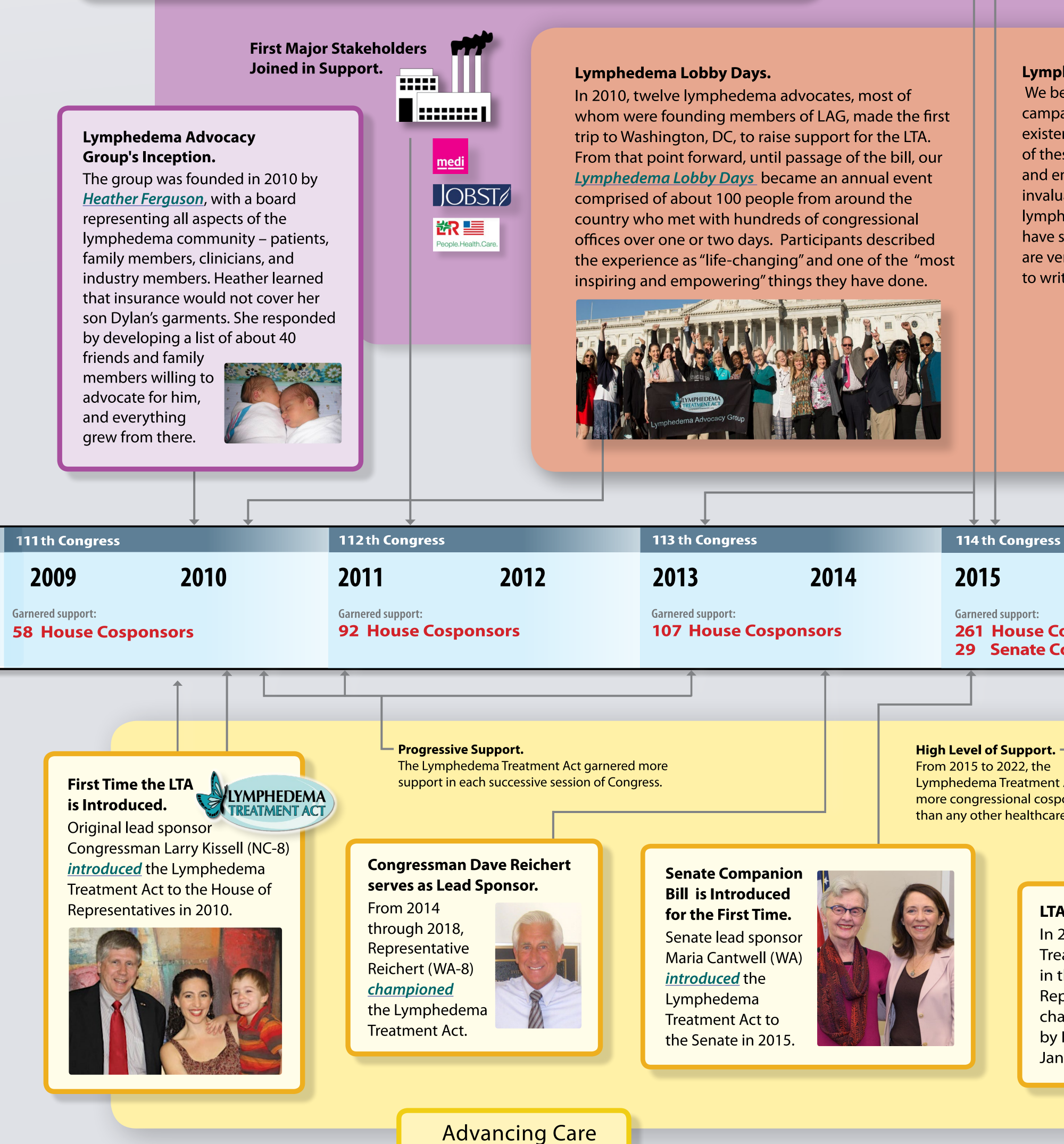


The Lymphedema Advocacy Group (LAG) is an all-volunteer patientcentered organization in the United States focused on improving access to care. Executive Director Heather Ferguson founded LAG in 2010, after the birth of her twins, one of whom has primary lymphedema.

This timeline depicts the group's 13-year journey of building a nationwide grassroots effort and learning to navigate the American legislative system. Key elements include networking, coalition building, training and activating

advocates to communicate with legislators, and promoting research findings that demonstrate a reduction in lymphedema-related healthcare spending through better patient outcomes.

LAG's successful strategies have resulted in wide-spread improvements to insurance coverage for compression garments and supplies for millions of Americans. We are happy to share our timeline as a blueprint for community-building, giving voice, and advancing care. Learn more about us at LymphedemaAdvocacyGroup.org.



State Teams.

We created regional *teams* of advocates in 2013, and by 2015 we had teams in all 50 states.

Advocacy Training.

thousands of congressional meetings.



Lymphedema Stories.

We began the <u>My Lymphedema Story</u> campaign during our first year of existence and have amassed over 1,000 of these deeply personal and often raw and emotional narratives. They are invaluable for helping others understand lymphedema's impact, and many patients have said their lymphedema journeys are very therapeutic

to write about.

TREATMENT ACT Selected Stories Stories and pictures from citizens across United States of America selected from our collec #400 letters sent to the Lymphedema Advocacy (g their experience with lymphedema and the imp passing the Lymphedema Treatment Act.

113 tl	n Cong	jress

2015

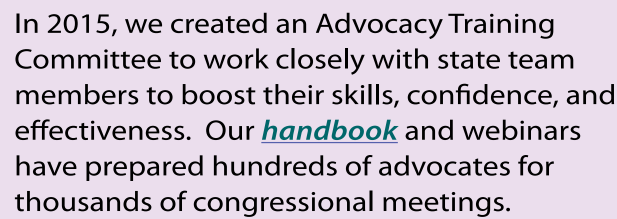
Garnered support: **261 House Cosponsors 29 Senate Cosponsors**

2016

ere S.	High Level of Support. From 2015 to 2022, the Lymphedema Treatment Act had more congressional cosponsors than any other healthcare bill.	Entire 100 member SNATE 3885 67 member SUPER MAJORITY MAJORITY MAJORITY MAJORITY MAJORITY	Sav In 2 con con
		SENATE HOUSE COSPONSORS 2022	pot
			ena

LTA Passes in the House. In 2019, the Lymphedema Treatment Act was *passed* in the House of Representatives, championed by House sponsor

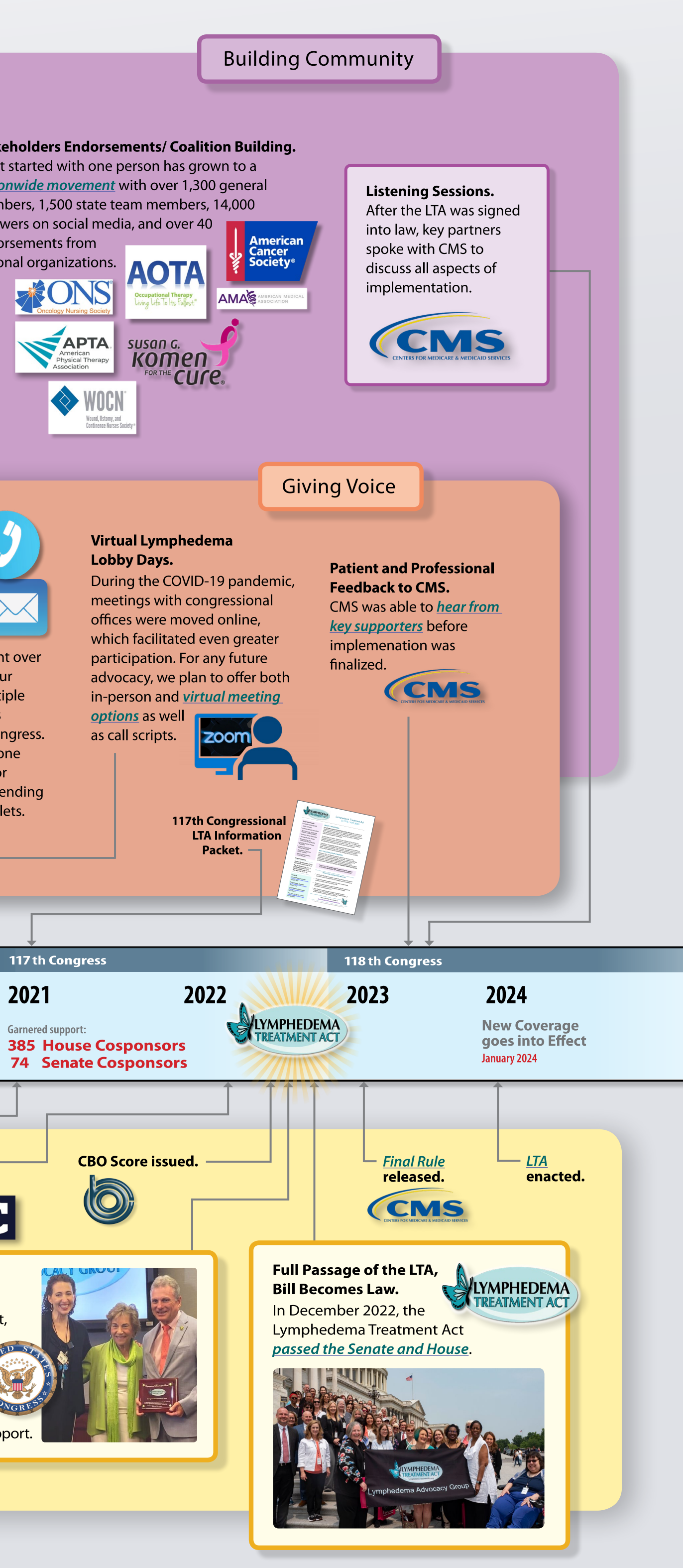
Jan Schakowsky (IL-9).











- *resources*, some geared toward the general information cards free of charge to doctors and therapists' offices, manufacturers and distributors, support groups, and anyone who wishes to help spread awareness. Two of our most popular full-page
- documents are "6 Things You May Not Know About Lymphedema" and
- "Lymphedema:



