Dear friends,

This past year we continued to see major leaps forward in the landscape of myeloma, with new treatments approved in Europe and a growing pipeline of forthcoming myeloma and AL amyloidosis medicines. Whilst these advances represent significant promise for the patient community in terms of options, outcomes and quality of life, further progress on access to medicines, education and supportive care needs and patient evidence gaps must accompany treatment innovations across Europe.

In this report, you’ll read about the key accomplishments of Myeloma Patients Europe (MPE) from 2023, many of which focus on these unmet needs. These included regularly providing members, patients, and carers with accurate and timely information, working with European and national stakeholders to advocate for improved access, understanding and improving the role of patients in treatment and care decision-making, developing tools to help reduce diagnosis delays and much more.

In addition to our programmatic accomplishments, MPE’s membership continues to grow as well as our contributions in national and European forums. Now with 49 members, MPE remains a strong and effective network of organisations working tirelessly to support myeloma and AL amyloidosis patients and to advocate for continued improvements in care, treatment, access, research, education, and patient involvement in drug development and HTA. Our collective voice is critical to drive change and each year that voice grows stronger.

We want to thank our members, patients, carers and advocates, we recognize and deeply appreciate the work you do every day as well as your continued support of MPE. We also want to thank our Medical Advisory Committee, industry sponsors, and stakeholders for your ongoing dedication to MPE and the patient community.

We look forward to building on these achievements in the year ahead and continuing to work together to make a difference for patients.

Best wishes,

LISE-LOTT ERIKSSON
President

KATE MORGAN
Co-Chief Executive Officer

KATIE JOYNER
Co-Chief Executive Officer
MYELOMA PATIENTS EUROPE

Myeloma Patients Europe (MPE) is an umbrella organisation of myeloma and AL amyloidosis patient groups across Europe. The mission of the organisation is to provide education, information, and support to member groups and to advocate at European, national, and local levels for the best possible research and equal access to treatment and care. To achieve its aims, MPE works directly with members, healthcare professionals, reimbursement authorities, regulators, politicians, pharmaceutical companies, and the media to ensure all stakeholders are collaborating to improve patient outcomes and reduce inequalities across Europe.

MPE’s Strategic Objectives

In 2019, MPE’s board and membership developed and confirmed seven strategic objectives to guide our work programme through 2024. All of the activities of MPE support one or more of the following:

- Provide a strong voice for myeloma and AL amyloidosis patients at a European and international level.
- Strengthen members’ and individual advocates’ ability to advocate effectively.
- Secure a larger patient voice in myeloma and AL amyloidosis research.
- Accelerate progress in the treatment and care of myeloma and AL amyloidosis patients.
- Increase the number of patients who have their myeloma and/or AL amyloidosis diagnosed in a timely manner.
- Improve access to optimal treatment and care for all myeloma and AL amyloidosis patients in Europe.
- Strengthen MPE’s effectiveness, sustainability and capacity to take collective action.

Secure a larger patient voice in myeloma and AL amyloidosis research.

Looking ahead: In 2024, MPE will develop a new strategic plan. We look forward to working with our members, stakeholders, and the community to align on a vision for the next five years and develop a roadmap to achieve our shared goals. Stay tuned!
In 2023, MPE had 49 full and associate members representing 31 different countries. MPE continues to grow and support its members to create a network of highly effective, efficient and sustainable patient organisations across Europe.
Full Members

Armenia
- Armenian Hematology Association

Austria
- Multiples Myelom Selbsthilfe Österreich (MMSOe)
- Myelom- und Lymphomhilfe Österreich

Belgium
- Contactgroep Myeloom en Waldenström Patiënten Vlaanderen vzw (CMP vzw)
- MyMu Wallonie-Bruxelles (MyMu)

Croatia
- Udruga za podršku oboljelima od multiplog mijeloma (Mijelom CRO)
- Hrvatska udruga leukemija i limfomi (HULL)

Denmark
- Dansk Myelomatose Forening

Finland
- Suomen Syöpäpotilaat – Cancerpatienterna i Finland ry

France
- Association Française des Malades du Myélome Multiple (AF3M)

Germany
- AMM-Online
- Deutsche Leukämie- & Lymphom-Hilfe e.V. (DLH)
- Myelom-Gruppe RHEIN-MAIN – Leukaemiehilfe RHEIN-MAIN e.V. (LHRM)
- Myelom.Online

Iceland
- Perluvinir

Ireland
- Multiple Myeloma Ireland

Israel
- AMEN Israel
- Amyloidosis Israel

Latvia
- Onkologisko pacientu atbalsta biedriba “Dzīvības koks”

Lithuania
- Asociacija “Kraujas”

Macedonia
- BORKA
- HEMA

Netherlands
- Stichting Hematon
- Stichting Amyloidose Nederland (SAN)

Norway
- Blodkreft Foreningen

Poland
- Fundacja Carita – Żyć ze Szpiczakiem
- Polskie Stowarzyszenie Pomocy Chorym na Szpiczaka

Portugal
- Associação Portuguesa Contra a Leucemia (APCL)
- Associação Portuguesa Leucemias e Linfomas (APLL)

Romania
- Myeloma Euronet Romania (MER)
- SOS Mielom
Russia
- Society for Assistance to Patients with Oncohaematological Diseases “Mercy Bridge”

Serbia
- Udruženje obolelih od multiplog mijeloma

Slovakia
- Združenie pacientov s hematológickými malignitami
- Slovenská myelómová spolocnost

Slovenia
- Drustvo bolnikov z limfomom (DBL)
- Slovensko Združenje Bolnikov Z
- Limfomom In Levkemijo L&L
- Društvo bolnikov s krvnimi boleznimi Slovenije

Spain
- Asociación Española de Amyloidosis (AMIKILO)
- Agrupación Española Contra la Leucemia y Enfermedades de la Sangre (AELCLES)

Sweden
- Blodcancerförbundet Sweden

Switzerland
- Stiftung zur Förderung der Knochenmarktransplantation Schweiz
- MPS Myelom Patienten Schweiz

Turkey
- KANKO/BIRKAN

Associate Members

Czech Republic
- Klub pacientů mnohočetný myelom, z.s.

Bosnia and Herzegovina
- Bosnian Myeloma Patient Group

Portugal
- Centro de Histocompatibilidade do Norte (CHN)

Canada
- Myeloma Canada

This year, MPE welcomed two new member organisations: Myeloma Canada and Myelom.Online in Germany!

MPE extends a special thank you to our members and the myeloma and AL amyloidosis patient community. Your work is invaluable to the patients and families you support, and we remain inspired by your unwavering efforts to improve patients' lives.
**MPE BOARD**

The MPE Board is multidisciplinary and responsible for the strategy, governance, and accountability of the organisation. The Board is comprised of myeloma patients, caregivers, and advocates.

**MPE’s 2023 Board Members**

- Lise-lott Eriksson, President (Sweden)
- Ananda Plate, Vice President (Germany)
- Barbara Leonardi, Treasurer (Poland)
- Riikka-Leena Manninen, Secretary (Finland)
- Reidar Nordby, Board Member (Norway)
- Diane Loening-Martens, Board Member (Germany)
- Zvika Zilberman, Board Member (Israel)

*In 2023 MPE welcomed new Board Member, Reidar Nordby, from the Norwegian Blood Cancer Association*
MPE MEDICAL ADVISORY COMMITTEE

The MPE Medical Advisory Committee (MAC) provides evidence-based recommendations to MPE and members in support of advocacy initiatives, as well as updates on myeloma and AL amyloidosis disease, diagnostics, and treatments. It is composed of key opinion leaders representing myeloma and AL amyloidosis specialties, as well as MPE’s member countries.

2023 Medical Advisory Committee members:

- Prof. Heinz Ludwig (Austria)
- Dr. Vsevolod Potapenko (Russia)
- Prof. Roman Hájek (Czech Republic)
- Dr. Paolo Milani (Italy)
- Dr. Ulf-Henrik Mellqvist (Sweden)
- Dr. Charlotte Pawlyn (United Kingdom)
- Prof. Pieter Sonneveld (The Netherlands)
- Prof. Michel Delforge (Belgium)
- Dr. Anna Sureda (Spain)
- Dr. Laurent Garderet (France)
- Dr. Moshe Gatt (Israel)
- Prof. Oliver Karanfilski (Republic of North Macedonia)
- Dr. Valdas Pečeliūnas (Lithuania)
- Prof. Katja Weisel (Germany)
- Prof. Sonja Zweegman (Netherlands)
- Prof. Meral Beksac (Turkey)
- Prof. Efstathios Kastritis (Greece)
- Prof. Monique Minnema (The Netherlands)
- Dr. Magdalena Olszewska-Szopa (Poland)

MPE is grateful for the time and expertise the MAC provides to our organisation. On behalf of the patient community, thank you for your support!

“MPE’s Medical Advisory Committee facilitates a critical dialogue between clinicians and the patient community. We’re all working to improve patient outcomes, and by sharing what I see in the clinic and what MPE sees through their programmes, haematologists and advocates can better support patients.”

Dr. Moshe Gatt, Hadassah Medical Center, Israel
MPE is comprised of an international team of professionals dedicated to meeting the needs of our members and the patient community. Through their efforts, MPE successfully develops and implements programmes and projects that fulfil the organisation's mission and strategic objectives.

MPE STAFF

Kate Morgan, Co-Chief Executive Officer
Katie Joyner, Co-Chief Executive Officer

Medical Education and Scientific Engagement
Solène Clavreul, Head of Medical Education and Scientific Engagement
Morine Maguri, Project Officer
Thisari Dharmapiya, Medical and Scientific Officer

Patient Evidence
Eilidh Duncan, Head of Patient Research
Silene Ten Seldam, Research Assistant

Member and Patient Community Programmes
Linda Christopher, Project Manager

Access and Policy
Anne-Pierre Pickaert, Access to Medicines Secretariat

Communications
Ana Vallejo, Head of Communications
Patricia Matamoros, Communications Officer

Finance and Operations
Riika Lempiainen, Finance Officer
Valentina Christodoulidou, HR and Operations Officer
Through the collective work of our members, staff, and stakeholders, MPE continues to address the challenges in myeloma and AL amyloidosis and deliver tools, evidence and information to help improve patient outcomes, experience and quality of life. Below provides a summary of the major accomplishments from each workstream.
ACCESS, POLICY AND REGULATORY

To ensure all patients have access to the best treatment and care and to represent their interests at the European level.

MPE’s Access, Policy and Regulatory workstream provides tools and data for MPE and our members to navigate the access landscape and advocate for improved access to medicines, as well as work closely with regulators, reimbursement bodies and policymakers to influence decisions and initiatives that impact access across Europe. MPE’s major accomplishments in 2023 include:

- Completed a comprehensive update of the Myeloma Access Atlas, supporting members in their advocacy efforts to overcome variation in access to treatment and care in Europe.
- Delivered tailored coaching and support to members on access challenges.
- Presented data on inequalities in access between Western and Central and Eastern Europe at the European Society for Medical Oncology (ESMO), as well as several roundtable discussions and stakeholder events.
- Supported colleagues across the cancer community in developing atlases in other disease areas.
- Appointed to and served on the EU HTA Stakeholder Network of the European Commission, actively engaging in the implementation of the EU HTA Regulation.
Published and presented research results on the use of quality-of-life (QoL) instruments and patient-reported outcome measures (PROMs) reported in clinical trials and published literature in myeloma.

Provided training on national health technology assessment (HTA) and reimbursement processes and how patient organisations can participate and influence decision-making.

Delivered written submissions to the European Medicines Agency (EMA) in support of two new medicines and facilitated patient representation for three assessment processes.

Continued to advocate for the representation of patients in decision-making on medicines, including through MPE representation on the EMA Patients’ and Consumers’ Working Party (PCWP) and on the Steering Group for the Patient and Citizen’s Interest Group (PCIG) at HTAi.

“MPE has provided invaluable support for the Association of Cancer Patients in Finland (Suomen Syöpäpotilaat ry) on an access challenge relating to daratumumab. They listened, provided advice, spoke to relevant stakeholders, mentioned it at key congresses and even wrote a supporting letter for us to engage with the Finnish Society of Myeloma Doctors. We have additionally discussed a direction from the Ministry of Social Affairs and Health related to daratumumab. We really enjoyed working with them and a combined voice definitely makes us stronger!”

Emma Andersson, Association of Cancer Patients in Finland
MPE's Medical Education and Scientific Engagement department promotes understanding around medicine, clinical trials, and research to strengthen advocacy efforts and to ensure access to the latest information on myeloma and AL amyloidosis developments. MPE reached the following milestones this year:

- Beta launch of Myeloma and AL Amyloidosis Clinical Trial Navigator, a patient-friendly search tool of recruiting clinical trials in Europe, which will undergo exhaustive testing before the official launch in 2024.

- Published new AL amyloidosis Q&As on diagnosis and monitoring, living with AL amyloidosis, and the differences between AL amyloidosis and myeloma, as well as a new AL amyloidosis patient guide.

- Published new myeloma educational materials including factsheets on approved treatments, resources on minimal residual disease (MRD), Q&As and a webinar on CAR-T therapy, and an updated myeloma patient guide.

- Produced video interviews and webinars with key opinion leaders on the latest myeloma and AL amyloidosis data from the European Hematology Association (EHA) and American Society of Hematology (ASH) Annual Congresses.

- Published research on the disease burden on the quality of life for AL amyloidosis patients and on the challenges and impacts of delayed diagnosis in AL amyloidosis.

“European patients with myeloma can have easier access to clinical trials. The website can serve as a trusted source of accurate and up-to-date information on myeloma clinical trials in Europe. Patients and their families can rely on it to find information that is verified by a reputable organisation.”

Ariunsanaa Baatar, Sanna Foundation, Hungary
PATIENT EVIDENCE

To support informed, data-driven decision making through the generation of evidence on patient needs and preferences.

MPE’s Patient Evidence department generates robust data to support decision-making in myeloma and AL amyloidosis and to advocate with scientific, industry, regulatory and policy decision-makers for improved care and treatment for patients across Europe. Key achievements from 2023 include:

Launched pan-European survey on patients’ and carers’ perspectives on how and where treatments are administered to inform decision-making among industry and regulatory stakeholders as they consider how drugs should be developed and administered.

Published research on patient and haematologist perspectives on Minimal Residual Disease (MRD) and hosted an online workshop to discuss the results.

Published research findings on the needs of working age myeloma patients, in collaboration with the University of Stirling and MPE members in Poland, Germany and Czech Republic.

Provided input into EORTC quality of life working group projects exploring financial toxicity, interpretation guidelines, and reference values, and began contributing to a new project investigating different modes of patient recruitment.

“"I have participated in a few research projects run by the MPE Patient Evidence Team. The results shed new light on the myeloma community and their lives. By collaborating with European universities, and patient organisations, the Patient Evidence team demonstrates MPE’s academic and border-breaking capabilities."

Barbara Leonardi, Carita Foundation, Poland

Commenced a patient and haematologist survey on shared decision-making, in partnership with KU Leuven, to understand how to improve the processes for patient involvement in treatment and care decisions.
MEMBER AND PATIENT COMMUNITY PROGRAMMES

To build and empower a network of effective patient advocates and organisations across Europe and to provide a strong platform for the voice of myeloma and AL amyloidosis patients to be heard.

MPE’s Member and Patient Community workstream is comprised of several distinct programmes designed to strengthen members’ and patients’ ability to advocate effectively and to build a network of impactful patient organisations across Europe. This year, we accomplished several key objectives:

- Provided scholarships for 10 member organisations to implement projects to meet the needs of their local patient communities, which included initiatives on psychological support for newly diagnosed patients, generating data to use in HTA assessments, general practitioner awareness programmes and much more.

- Launched quarterly MPE Member meetings to highlight the ongoing achievements of national organisations and provide a forum to exchange ideas among members.

- Trained seven new participants through the MPE Advocate Development Programme, continuing our efforts to build a network of active and empowered patient advocates.

“The Advocate Development Programme (ADP) exceeded all my expectations. The program is very well-structured, with an incredible panel of speakers who were always willing to help us. I am deeply thankful to the entire MPE team for the knowledge, expertise, support and genuine care they provided me. Being part of the programme was pivotal because, since I joined the ADP, I have had and continue to have the chance to participate in different projects, with the aim of helping and empowering multiple myeloma patients.”

Ana Filipa Lopes, Portuguese Association Against Leukemia (APCL)
Hosted 80 members, industry partners, and other stakeholders at the MPE Masterclass with educational sessions addressing treatment updates, patient involvement in HTA, clinical trial access and the most pressing needs of the patient community.

“Nothing is more powerful than learning from each other, and that is exemplified in the MPE Masterclass. For the past two years, I have had the pleasure of presenting at the Masterclass and working together with the MPE members on patient and advocate involvement in HTA. I have been struck by the level of knowledge, commitment and sheer power of the MPE community to make change happen. Through training sessions addressing the key needs of patient advocates, the Masterclass empowers the community to work together to address some of the most challenging healthcare issues.”

Neil Bertelsen, Steering Committee Member and past Chair of HTAi PCIG

Held European Myeloma Day on September 27, 2023, uniting patients, carers, advocates, clinicians, medical societies, industry and other stakeholders to raise awareness of myeloma and the need for a timely diagnosis for every patient.
Towards a Timely Diagnosis for All Patients

Increase the number of patients who have their myeloma and/or AL amyloidosis diagnosed in a timely manner.

To raise awareness about myeloma and AL amyloidosis, and help reduce delays in diagnosis for each disease, MPE developed the European Myeloma Diagnosis Pathway and the European Amyloidosis Diagnosis Pathway. These tools are designed to help general practitioners (GPs) recognize the signs and symptoms earlier and which tests to undertake if myeloma or amyloidosis is suspected.

To ensure wide-spread dissemination, MPE and our members translated the myeloma pathway into 7 languages and member organisations led national efforts to disseminate the tool to GPs through congress attendance, info booths at GP events and direct outreach.

The amyloidosis pathway was presented at the Spanish Hematology Congress and will be translated into additional languages in the year ahead.

Over the next year MPE will build on this momentum and reach hundreds, if not thousands, of GPs across Europe with the pathways.
Sharing the myeloma diagnosis pathway had a great impact in local communities in raising awareness about myeloma. We distributed 4,000 copies of the pathway and educational materials about multiple myeloma to health centres throughout Serbia. Every day we receive pictures of our members from all around Serbia from local clinics, where we can see how posters and flyers are distributed in smaller communities and villages throughout the country. Due to poor knowledge about the symptoms of myeloma among most GPs, we have received invitations from some local clinicians to organize lectures on the topic of myeloma together with leading haematologists.”

Snezana Doder, Association of Myeloma Patients Serbia
At the onset of the crisis in Ukraine, MPE worked quickly to understand the immediate impact of the war on cancer patients, including myeloma and AL amyloidosis patients in Ukraine or fleeing from Ukraine. As the war continues, MPE remains committed to ensuring that patients and carers receive the help they need in accessing treatment and care during this crisis.

Throughout 2023, MPE assisted in the translation of medical records to help expedite treatment and linked patients who have fled Ukraine and need myeloma, AL amyloidosis, haematology, or other cancer support in new countries. We will continue to provide support and services for as long as necessary.
Through presentations at congresses and meetings, MPE contributes to important discussions about myeloma treatment and care, patient advocacy and patient evidence. Some key highlights from 2023 are:

- A presentation at the EHA annual congress by Kate Morgan, MPE Co-Chief Executive Officer, on the importance of measuring quality of life in myeloma clinical trials during an EHA Guidelines Session.

- A poster presentation at the International Myeloma Society (IMS) annual meeting by Silene ten Seldam, MPE Research Assistant, on patient and clinician perspectives on minimal residual disease.

- A presentation at the European Society for Medical Oncology (ESMO) 2023 given by Kate Morgan on “A gap analysis on access to modern medicines and treatments: A comparison between West and Eastern Europe.”

- An ISPOR 2023 poster presentation, in collaboration with KU Leuven, of work supported by MPE titled “Needs and Experiences of Patients and Healthcare Professionals Towards Shared Decision Making in Multiple Myeloma Clinical Practice: A Qualitative Study in Europe.”

- A presentation on “AL amyloidosis diagnosis: challenges and recommendations” given by Ana Vallejo, MPE Head of Communications, during a World Amyloidosis Day event organised by the Spanish Amyloidosis Association (AMILO), in collaboration with the Spanish Society of Hematology.

- Solene Clavreul, MPE Head of Medical Education and Scientific Engagement, co-authored a poster at the ASH annual meeting entitled "Cytokine Release Syndrome: The Patient, Caregiver and Healthcare Professional Experience."

- Presentation at the European Myeloma Network (EMN) annual meeting of MPE’s Myeloma Access Atlas, given by Kate Morgan.

- Hans Scheurer, former President of MPE, presented the patient perspective during a round table on “Ethical issues in CAR T-cell treatment” hosted by the European School of Hematology.
EUROPEAN COMMISSION PROJECTS

To provide a strong voice for myeloma and AL amyloidosis patients at European and international levels and collaborate with key stakeholders to take collective action on behalf of the patient community.

MPE is involved in several European projects, ensuring that the perspectives of the myeloma, AL amyloidosis, and the broader haematology and oncology patient communities are represented, and that deliverables are designed with patients’ needs in mind. Key accomplishments for each project in 2023 include:

SISAQOL-IMI
SISAQOL-IMI (Setting International Standards of Patient-Reported Outcomes and Quality of Life Endpoints in Cancer Clinical Trials – Innovative Medicines Initiative), is an international consortium convened to generate recommendations to standardise patient reported outcome (PRO) data in cancer clinical trials.

- Held in-person workshop at the WECAN Academy in Frankfurt, Germany educating patient advocates on the challenges in designing, analysing and reporting PRO data and the importance of SISAQOL-IMI to standardise the way this is done.
- Facilitated input of patient advocates across oncology into the plain language glossary, which aims to allow all stakeholders, including patients and patient organisations, to have a better understanding of the context and terminology used in PRO data collection, analysis and reporting.
- Developed plain language executive summaries to ensure transparency for the patient community of the project’s process.

"MPE, representing WECAN, is an invaluable contributor to SISAQOL-IMI. From involving patients in each work package to organising patient meetings with work package leaders, MPE is ensuring the patient voice is heard and incorporated throughout the project and its deliverables. SISAQOL-IMI is a technical project involving setting standards on the design, analysis and interpretation of quality of life and patient reported outcome data. MPE exemplifies a way of working that bridges the communication gap between the experts and the patient community. The commitment and openness of MPE to make sure that the results are truly relevant for the patients has made SISAQOL-IMI even more meaningful to all project partners and different stakeholders."

Madeline Pe, PhD, Head of Quality of Life Department, EORTC
HORIZON EUROPE ASCERTAIN
is a four-year project consortium, led by the University of Erasmus MC in Rotterdam, exploring alternative models of pricing, cost-effectiveness and reimbursement to improve access to innovative therapies for patients.

HORIZON2020 CARAMBA
is a five-year research collaboration between ten partners from six EU-countries exploring Chimeric Antigen Receptor T-cell therapy (CAR-T) for the treatment of myeloma.

IMI HARMONY
is a consortium of 50 haematological partners, aiming to facilitate personalised medicine, identify the most effective treatments and improve diagnosis by collecting and disseminating valuable clinical data on haematologic malignancies through a big data platform.

- Developed an iterative patient engagement plan to ensure the patient perspective is included across all work packages and deliverables.
- Provided patient input in all work package meetings and across deliverables to-date.
- Planned educational resources and focus groups on drug pricing for patient advocates.

- Completed research on patient evidence requirements in the regulatory and reimbursement assessment of CAR-T cell therapies, and on the patient experience and PRO collection in CAR-T clinical trials.
- Developed and published educational materials (webinars, Q&As, interviews etc.) on CAR-T, focusing on the patient experience, clinician perspective, side effects and manufacturing.
- Provided patient perspectives on the CARAMBA trial results and next steps.

- Provided input into research proposals and analyses to ensure the patient voice is represented in all studies.
- Advocated for the addition of PRO and QoL data to the HARMONY platform, and for better quality data.
- Provided feedback on HARMONY sustainability plans from a patient perspective.
MPE 2023 FUNDERS

Pharmaceutical industry supporters:

- AbbVie
- Amgen
- Alexion AstraZeneca
- BeiGene
- Binding Site
- Bristol Myers Squibb
- GlaxoSmithKline
- Janssen
- Menarini Stemline
- Novartis
- Oncopeptides
- Pfizer
- Prothena Biosciences
- Regeneron
- Sandoz
- Sanofi
- Sebia
- SkylineDx
- Roche
- Takeda

Public supporters:

- Horizon 2020 ASCERTAIN
- Horizon 2020 CARAMBA
- IMI HARMONY
- SISAQOL IMI

Thank you!

MPE is very grateful for the continued commitment of our funders and would like to thank them for their support this year.

MPE works with key stakeholders to strengthen the voice of myeloma and AL amyloidosis patients, advocate for the patient community’s needs and interests and work collectively to improve patient outcomes and experiences across Europe. Thank you for your continued partnership:

COMMUNITY COLLABORATION

- ECO
- EHA
- EMA
- EMN
- EORTC
- ESMO
- EU HTA Stakeholder Network
- EURORDIS
- HTAI
- PFMD
- WECAN
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