Oral Abstracts and Posters

	MONDAY MAY 27, 2024 ORAL ABSTRACTS	AND POSTERS
Poster ID (submission ID)	DIAGNOSIS OF AMYLOID	
OA1 (#496)	A feasibility study into computer vision quantification of cardiac amyloid burden in endomyocardial biopsies	Taxiarchis Kourelis Mayo Clinic
OA2 (#99)	AmyLite Assay Quantifies Kinetically Unstable Circulating	Xin Jiang
	Amyloidogenic Lambda FLC – Diagnostic and Prognostic Implications for Lambda AL Amyloidosis	Protego Biopharma
OA3 (#420)	Lumbar spinal stenosis (LSS) and carpal tunnel syndrome (CTS) as surrogates for wild-type transthyretin amyloid cardiomyopathy (ATTR-CM)	Laura De Michieli Universita degli Studi di Padova
OA4 (#564)	Screening for Amyloidosis at the time of Carpal Tunnel Release Surgery in Real World Practice is a Successful Strategy at Diagnosing Early Cardiac Amyloidosis	Mazen Hanna Cleveland Clinic
	BASIC SCIENCE AL AMYLOIDOSIS	
OA5 (#464)	Bone marrow-free sequencing of M protein genes: a liquid biopsy approach in monoclonal gammopathies	Alice Nevone Regione Lombardia
OA6 (#253)	Characterization of the Peptide-Antibody Fusion, AT-02 – Studies to Support its Use as an Immunotherapy in Patients with Amyloidosis	Jonathan Wall University of Tennessee
OA7 (#38)	Helical superstructures between amyloid and collagen VI in heart-derived fibrils from a patient with Light Chain Amyloidosis.	Stefano Ricagno Universita degli Studi di Milano
OA8 (#181)	Patient derived AL amyloid induces cellular toxicity in macrophages, hepatocytes, and cardiomyocytes upon cellular engulfment of amyloid material	Joseph Jackson The University of Tennessee System
OA9 (#333)	Single-cell, spatial analysis of the renal AL immunome supports a T-cell mediated tissue toxicity mechanism.	Charalampos Charalampous Mayo Clinic
	AL AMYLOIDOSIS PROGNROSIS AND TREATMENT	
OA10 (#466)	A phase II study of daratumumab and pomalidomide in previously treated patients with AL amyloidosis	Paolo Milani Universita degli Studi di Pavia
OA11 (#209)	Clinical Factors Associated with Early Sudden Unexpected Death in Systemic AL Amyloidosis: Insights from 138 Cases	Andrew Staron Boston Medical Center
OA12 (#267)	An ECG-Echo Risk Score for Systemic Light Chain Amyloidosis	Aparna Hari Mata Amritanandamayi Math
OA13 (#529)	Daratumumab-Based Front-line Therapy Improves Treatment Response and Survival in Patients with Immunoglobulin Light chain (AL) Amyloidosis: The Mayo Clinic Experience	Binoy Yohannan Mayo Clinic
OA14 (#185)	Efficacy and Safety of Daratumumab Monotherapy in Newly Diagnosed Patients with Stage 3B Light-Chain Amyloidosis: A Phase 2 Study by the European Myeloma Network	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon



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OA15 (#273)	Ethnicity in AL amyloidosis shows underestimation of disease risks by biomarker-based staging for ethnic minorities patients in systemic AL amyloidosis	Jahanzaib Khwaja HM Government of the UK of Great Britain and Northern Ireland
OA16 (#501)	Outcomes of patients with AL amyloidosis and end-stage renal disease after initiation of therapy	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon
OA17 (#264)	Peripheral Neuropathy in AL Amyloidosis: Clinical Presentations and Outcomes of Current Therapies	Pitcha Chompoopong Regents of the University of Minnesota
OA18 (#235)	Plasma cell characteristics predict benefit from intensified therapy in AL amyloidosis	Maximilian Steinhardt Julius-Maximilians-Universitat Wurzburg
OA19 (#475)	Proposed hematologic progression criterion in patients with AL amyloidosis.	Giovanni Palladini Universita degli Studi di Pavia
	AI/ELECTRONIC RECORDS TO FACILITATE DIAGNOSIS	
PA1 (#12)	A Qualitative Study of Artificial Intelligence-Based Tools to Raise Suspicion for ATTR Cardiomyopathy	Baljash Cheema Northwestern University
PA2 (#69)	Machine Learning-Based Clustering Identifies Novel Subgroups of Patients with AL Amyloidosis with Distinct Clinical Characteristics	Shankara Anand Boston University
PA3 (#105)	Artificial intelligence for the detection of systemic amyloidosis	Delfina Cirelli Hospital Italiano de Buenos Aires
PA4 (#122)	Sex differences in the performance of the transthyretin amyloid cardiomyopathy (ATTR-CM) risk score	M Trejeeve Martyn Cleveland Clinic Foundation
PA5 (#147)	Timing and co-occurrence of red-flag diagnoses prior to a diagnosis of systemic light chain (AL) amyloidosis	Anita D'souza Medical College of Wisconsin
PA6 (#164)	Raising awareness to support early detection and diagnosis of Transthyretin (TTR) Amyloidosis in Ghana/West Africa from a public and health professionals trainers and trainees perspective	Kwaku Appiah-Kubi C K Tedam University of Technology and Applied Sciences
PA7 (#198)	Characterization of ATTR Amyloidosis by Phenotype from Claims Data in the United States and Japan: Preliminary Results from the OverTTuRe Study	Kevin Alexander Stanford University
PA8 (#271)	Improving Detection of AL Amyloidosis (IDEA)—testing Al cardiac algorithms in the real world	Angela Dispenzieri Mayo Clinic
PA9 (#366)	Prometeo: a simple and accurate amyloidosis subtyping by MS-based proteomics.	Pierluigi Mauri Consiglio Nazionale delle Ricerche
PA10 (#457)	Hereditary Transthyretin-related Amyloidosis ongoing clinical study: a baseline report of the first 3,167 participants	Pierre Engel Centogene GmbH

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PA12 (#516)	The value of AI for enhancing suspicion of cardiac amyloidosis using electrocardiography and echocardiography: A narrative review	Martha Grogan Mayo Clinic h
PA13 (#544)	Evolving knowledge of "red flag" clinical features associated with TTR p.Val142lle in a diverse electronic health record-linked biobank	Amy Kontorovich Mount Sinai Health System
PA14 (#545)	Machine Learning to Predict Mortality among Patients with Transthyretin Amyloid Cardiomyopathy	Ruizhi Liao Empallo Inc
PA15 (#551)	Measuring ATTRv-neuropathy in real world practice: a proposed protocol	Wilson Marques Junior Governo do Estado de Sao Paulo
PA16 (#561)	Exploring Patient- and Provider Characteristics Associated With The Utilization of Artificial-Intelligence-based Models to Detect Cardiac Amyloidosis (CA): A Subset Analysis From the On-going PREDICT-AMY Trial	Eli Muchtar Mayo Clinic
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PA18 (#32)	The Cryo-EM structure of renal amyloid fibrils suggests structurally homogeneous multiorgan aggregation in AL amyloidosis.	Sarita Sarita Universita degli Studi di Milano
PA19 (#37)	Gene Expression Sets and Renal Profiling from the RAIN (Renal AL Amyloid Involvement and NEOD00) Trial	Cindy Varga Atrium Health
PA20 (#39)	Determinants of amyloidogenic behavior in AL amyloidosis patient-derived AL55 light chain: Insights from structural and biophysical studies	Sarita Sarita Universita degli Studi di Milano
PA21 (#40)	The Cryo-EM structure of renal amyloid fibrils suggests structurally homogeneous multiorgan aggregation in AL amyloidosis	Sarita Sarita Universita degli Studi di Milano
PA22 (#54)	Bone Marrow Interstitial Amyloid and Its Microenvironment	Ping Zhou Tufts University
PA23 (#59)	Intact IgG in Light Chain Amyloidosis	Olga Gursky Boston University
PA24 (#64)	Recombinant light chain production and analyses for the development of genetic diagnostic tests for AL	Huyen Phan Westmead Institute for Medical Research
PA25 (#70)	Role of the C-terminus disulfide bond in amyloid fibril formation of full-length human immunoglobulin $\lambda 6a$ and Wil light chains	Lindsey Lampe Mayo Clinic
PA26 (#117)	Incidence of transthyretin amyloid cardiomyopathy from a French nationwide study of in- and out-patient databases	Thibaud Damy Republique Francaise

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PA28 (#140)	Enhanced stabilisation of an amyloidogenic light chain using a tight binding variable heavy domain to mimic the homodimer complex	Alana Maerivoet University of Liverpool
PA29 (#168)	Deciphering the conformational landscape of amyloidogenic lambda light chains associated with AL amyloidosis	Sarita Sarita Universita degli Studi di Milano
PA30 (#170)	Amyloid free light chains disturb calcium transients and contractility in an AL cardiac amyloidosis 3D human spheroid model	Mélanie Bézard Republique Francaise
PA31 (#188)	Analyzing the contribution of neutrophils in amyloid clearance using murine amyloidoma models	Trevor Hancock University of Tennessee System
PA32 (#207)	Developing Conformation-Sensitive Antibodies Targeting Amyloid Aggregates of Lambda-6 Light Chains: A Structure-Based Approach for Therapeutic Intervention	Luis Del Pozo-Yauner University of South Alabama
PA33 (#210)	A SNAP23-mediated SNARE complex is necessary for Ig free light chain secretion in AL amyloidosis and Multiple Myeloma, representing a novel molecular target	Emre Karayol Harvard University
PA34 (#215)	Heterohybridomas Producing Human Light Chains: Success with CD138+ Cells from Myeloma and Polyclonal Gammopathy But Not AL Patients	Ping Zhou Tufts University
PA35 (#222)	AL-Base: An updated resource for analyzing amyloidogenic antibody light chain sequences	Gareth Morgan Boston University
PA36 (#254)	Characterization of a Novel Beta-sheet Peptide-Fc Fusion for Targeting Systemic Amyloid Deposits	Jonathan Wall University of Tennessee System
PA37 (#259)	Sequencing of amyloidogenic monoclonal immunoglobulin light chain repertoires from AL patients by combing RNA-based assay and mass spectrometry	Shuang Wang Peking University
PA38 (#269)	Non-coding RNAs regulate novel signaling pathways in AL amyloidosis that are targetable by FDA approved drugs	Oshrat Hershkovitz-Rokah Maccabi Healthcare Services
PA39 (#278)	Cardiotoxicity in light chain amyloidosis: Insights from a murine model with intramyocardial injection of patient-derived amyloidogenic light chains.	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon
PA40 (#294)	Temporal Changes in the Renal Cytokine Profile in Response to AA Amyloidosis Induce Macrophage Infiltration Enabling Host-Mediated Targeting of Therapeutic Chimeric Antigen Receptor Macrophages (CARM)	Manasi Balachandran The University of Tennessee System
PA41 (#338)	Truncation of the constant domain drives amyloid formation by immunoglobulin light chains: towards a physiological fibrillogenesis model	Francesca Lavatelli University of Pavia
PA42 (#391)	Kinetic evidence for multiple aggregation pathways in antibody light chain variable domains	Sherry Wong Boston University

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PA45 (#463)	N-glycosylation of clonal immunoglobulin light chains as a risk factor for AL amyloidosis: Benchmarking N-glycosylation prediction tools	Alice Nevone Regione Lombardia
PA46 (#517)	RNA-based Immunoglobulin repertoire sequencing is a useful tool for prediction and management of AL amyloidosis along with monoclonal gammopathies of clinical significance	Murielle ROUSSEL Universite de Limoges
PA47 (#522)	Characterization of cardiac AL amyloidosis in a transgenic mouse model	Christophe Sirac Universite de Limoges
PA48 (#525)	Single-Cell Mass Cytometry Analysis Reveals a Prominent Immune Suppressive Signature in AL Amyloidosis	Theophilus Tandoh City of Hope
PA49 (#560)	Hemostasis dysfunction induces senile APOA2 amyloidosis in a mouse model	Christophe Sirac Universite de Limoges
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PA52 (#49)	V122I TTR and AL κ-type in a Patient with Cardiac Involvement and Spinal Stenosis	Xia Wu Tufts University
PA53 (#61)	Positive cardiac scintigraphy with Tc-99m DPD-SPECT: Diagnosis is not always ATTR cardiac amyloidosis	loannis Panagiotopoulos Onaseio Kardiocheirourgiko Kentro
PA54 (#65)	Prevalence, Incidence, And Characterization Of Light Chain Amyloidosis In The Usa: A Real-World Analysis Utilizing Electronic Health Records (EHR)	Pedro Laires AstraZeneca PLC
PA55 (#87)	Increasing Prevalence and Incidence of AL Amyloidosis Among Older Adults in the US	Preeti S. Bajaj, Phd Prothena Biosciences Inc.
PA56 (#119)	Transverse Carpal Ligament Analysis as Screening Tool for Amyloidosis	Seunghyuk Daniel Yang Trinity Health
PA57 (#124)	Predictors factors of mortality in advanced cardiac AL Amyloidosis:A prospective Cohort study - for improving cardiac stratification	Amira Zaroui Universite Paris-Est Creteil Val de Marne
PA58 (#177)	A Subset Of Patients With Renal AL-Amyloidosis Present Without Significant Proteinuria And Display Poor Renal Outcomes	Camille Cohen Republique Francaise
PA59 (#179)	The incidence of AL amyloidosis in patients with carpal tunnel syndrome	Gaja Cvejić Vidali Univerza v Ljubljani

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PA61 (#249)	There is more than meets the eye: an unexpected case of cardiac Amyloidosis.	Andrea Nevi Universita degli Studi di Verona
PA62 (#255)	Renal AL amyloidosis : an unusual presentation	Tarek Ashour Cleveland Clinic
PA63 (#308)	Role of Amyloidosis screening in a Monoclonal Gammopathy of Undetermined Significance (MGUS) clinic: Results from a pilot study at the University of Calgary	Victor Jimenez Zepeda Alberta Government
PA64 (#309)	Referral patterns in AL and ATTR Amyloidosis: Initial experience from the Amyloidosis Program of Calgary: Role of a Multidisciplanary approach	Victor Jimenez Zepeda Alberta Government
PA65 (#311)	Clinical significance of the Monoclonal Gammopathy of Undetermined Significance (MGUS) -like phenotype in patients with AL amyloidosis treated with Bortezomib-Containing Regimens (BCR) at the Amyloidosis Program of Calgary (APC)	Victor Jimenez Zepeda Alberta Government
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PA67 (#347)	Tracheobronchial Amyloidosis is Composed of Polytypic Immunoglobulins: A Report of 603 Cases	Daniel Larson Mayo Clinic
PA68 (#360)	Amyloid typing by liquid chromatography-tandem mass spectrometry analysis of unfractioned unfixed abdominal fat aspirates	Giulia Mazzini Regione Lombardia
PA69 (#363)	Latvian National Amyloidosis Registry Data 2020-2023	Valdis Ģībietis Rigas Stradina Universitate
PA70 (#368)	To Treat or Not to Treat – a Rare Case of Cardiogenic Shock and Isolated Cardiac Amyloidosis	Annie Mcgregor Rush University
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PA73 (#407)	An Unusual Journey: From Systemic to Localized AL Amyloidosis	Roberta Shcolnik Szor Hospital 9 de Julho
PA74 (#410)	Challenges in Amyloidosis Typing: The Role of Mass Spectrometry in a Middle-Income Country	Roberta Shcolnik Szor Hospital 9 de Julho
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PA76 (#419)	Subacute Axonal Polyradiculoneuropathy Presentation in AL and TTRv Amyloidosis	David Polston Cleveland Clinic

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PA82 (#486)	Pulmonary and nodal light chain amyloidosis as a presenting feature of Waldenstrom's Macroglobulinemia: A case report	Aaron Lobo Yale New Haven Health System
PA83 (#503)	De Novo AL Amyloidosis In Renal Allograft-Case Report And Literature Review	Batuhan Bulan Istanbul Universitesi-Cerrahpasa
PA84 (#520)	Cardiac Amyloidosis and Cardiac Sarcoidosis in a Patient with Heart Failure	Jennifer Maning Northwestern Memorial HealthCare
PA85 (#534)	Recurrent pleural effusion and AL amyloidosis diagnosis	Diane Xavier De Ávila Fluminense Federal University
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	PATHOLOGY	Truminense i ederal Omversity
PA86 (#55)	PATHOLOGY Clinical and genetic profiles of patients with hereditary and wild-type transthyretin amyloidosis: the Transthyretin Cardiac Amyloidosis Registry in the state of São Paulo, Brazil (REACT-SP)	Fabio Femandes Governo do Estado de Sao Paulo
PA86 (#55) PA87 (#57)	Clinical and genetic profiles of patients with hereditary and wild-type transthyretin amyloidosis: the Transthyretin Cardiac Amyloidosis Registry in the state of São Paulo,	Fabio Femandes Governo do Estado de Sao
	Clinical and genetic profiles of patients with hereditary and wild-type transthyretin amyloidosis: the Transthyretin Cardiac Amyloidosis Registry in the state of São Paulo, Brazil (REACT-SP) Pathological features in patients with hereditary	Fabio Femandes Governo do Estado de Sao Paulo Tomoaki Taguchi
PA87 (#57)	Clinical and genetic profiles of patients with hereditary and wild-type transthyretin amyloidosis: the Transthyretin Cardiac Amyloidosis Registry in the state of São Paulo, Brazil (REACT-SP) Pathological features in patients with hereditary transthyretin amyloidosis long after liver transplantation. Global pattems of tissue typing in amyloidosis: Results of	Fabio Femandes Governo do Estado de Sao Paulo Tomoaki Taguchi Kumamoto Daigaku Hironobu Naiki
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PA87 (#57) PA88 (#58) PA89 (#67) PA90 (#79)	Clinical and genetic profiles of patients with hereditary and wild-type transthyretin amyloidosis: the Transthyretin Cardiac Amyloidosis Registry in the state of São Paulo, Brazil (REACT-SP) Pathological features in patients with hereditary transthyretin amyloidosis long after liver transplantation. Global pattems of tissue typing in amyloidosis: Results of a survey by the International Society of Amyloidosis (ISA) Double Trouble: A case of biopsy proven ATTR and AL cardiac amyloid Unraveling the protection of neuronal cytoskeleton and synaptic structures by TTR Gallbladder Amyloid is Often Unexpected and May Have	Fabio Femandes Governo do Estado de Sao Paulo Tomoaki Taguchi Kumamoto Daigaku Hironobu Naiki Fukui Daigaku Arianne Clare Agdamag Cleveland Clinic Isabel Cardoso Universidade do Porto Catherine Hagen
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PA96 (#205)	Amyloid Infiltration of the Skeletal Muscle Infiltration is Common With Cardiac Amyloidosis	Sarah Cuddy Harvard University
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PA98 (#372)	The Detection Yield of Surrogate Tissue Biopsies across Classes of Systemic Amyloidosis: Review of 4,027 Cases	Natasha Burke Boston Medical Center
PA99 (#377)	Salivary Gland Amyloidosis: Proteomic Identification and Clinicopathologic Characterization of 57 cases.	April Chiu Mayo Clinic
PA100 (#413)	Quantitation of daratumumab among AL amyloidosis patients by M-quant measurement	Angela Dispenzieri Mayo Clinic
PA101 (#422)	Gastrointestinal Amyloid Screening Study (GASS): Is Screening for Amyloid in the Gastrointestinal Tract Useful?	Rola Khedraki Scripps Health
PA102 (#469)	Healthcare Amyloidosis European Registry (HEAR REGISTRY): Study design and methods	Mounira Kharoubi Universite Paris-Est Creteil Val de Marne
PA103 (#479)	Comparison of two free light chain assays: performance of the free light chain ratio as risk factor for MGUS progression	Qian Wang Mayo Clinic
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PA108 (#95)	Incidence of Second Primary Malignancies in Patients with AL amyloidosis and The Impact of Disease Stage and Therapies	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon
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PA110 (#108)	Epidemiological perspectives of amyloidosis in Argentina: results of an analysis of incidence and mortality in a population affiliated to a Medical Care Program	Delfina Cirelli Hospital Italiano de Buenos Aires
PA111 (#131)	Clinical Features of AL Amyloidosis Patients Harboring Clonal Hematopoiesis of Indeterminate Potential	Paolo Lopedote Steward Health Care

PA112 (#143)	Thrombotic and Bleeding complications in patients with AL Amyloidosis	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon
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PA114 (#160)	Is the Cardiac Amyloidosis Artificial Intelligence Electrocardiography (CA-AI-ECG) model useful in predicting outcomes in Multiple Myeloma (MM) patients without known Amyloidosis undergoing Autologous Stem Cell Transplant (ASCT)?	Angela Dispenzieri Mayo Clinic
PA115 (#186)	Identifying Early Suboptimal Hematological Response in Patients with AL Amyloidosis Treated with Bortezomibbased Therapy.	Nirija Ranjit Anderson Queensland Government
PA116 (#209)	Clinical Factors Associated with Early Sudden Unexpected Death in Systemic AL Amyloidosis: Insights from 138 Cases	Andrew Staron Boston Medical Center
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PA118 (#280)	Validating the performance of renal staging in AL amyloidosis patients undergoing autologous stem cell transplantation	Eli Muchtar Mayo Clinic
PA119 (#299)	Clinical profile and treatment outcomes in primary (AL) amyloidosis from low- and middle-income country (LMIC)	Pankaj Malhotra Government of India
PA120 (#332)	Mortality patterns among patients with cardiac amyloidosis in a tertiary care center in Latin America	Javier Torres Estado Peruano
PA121 (#348)	Predictive value of free light chain burden in patients newly diagnosed with AL amyloidosis treated with CyBorD or DaraCyBorD	Brendan Saunders Harvard University
PA122 (#354)	Inclusion criteria of clinical trials select patients with AL amyloidosis with favorable outcome and exclude almost one half of the real-life population.	Claudia Bellofiore Universita degli Studi di Pavia
PA123 (#355)	Long-term evaluation of amyloidosis diseases in Germany: National Clinical Amyloidosis Registry	Ute Hegenbart Universitat Heidelberg
PA124 (#361)	Predictors of timely and deep renal responses in AL amyloidosis	Matthew Rees Mayo Clinic
PA125 (#367)	Predictors of timely and deep cardiac responses in AL amyloidosis	Matthew Rees Mayo Clinic
PA126 (#384)	Refining prognostication in systemic AL amyloidosis	Jahanzaib Khwaja HM Government of the UK of Great Britain and Northern Ireland
PA127 (#392)	Long-term hematologic remission in Immunoglobulin Light Chain (AL) Amyloidosis	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon

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PA129 (#456)	A predictive model for day-100 transplant-related mortality in AL amyloidosis	Eli Muchtar Mayo Clinic
PA130 (#467)	IgM Immunoglobulin light chain amyloidosis: the quest for prognostic biomarkers in a rare disease with two distinct clonal phenotypes	Marco Basset Regione Lombardia
PA131 (#468)	Outcomes of patients with newly diagnosed light chain amyloidosis according to eligibility for clinical trials: Experience of a single institution	Jose Miguel Mateos Perez Hospital Clinic de Barcelona
PA132 (#490)	A EUropean REgistry and sample sharing networK to promote the diagnosis and management of light chain Amyloidosis: The EUREKA study	Mario Nuvolone Universita degli Studi di Pavia
PA133 (#491)	Real-life experience on light chain cardiac amyloidosis: delay diagnosis is still a major issue	Morgane Thiry Universite catholique de Louvain
PA134 (#494)	Clinical implications of genetic interphase fluorescence in situ hybridization aberrations in systemic light chain amyloidosis	Sara Oubari Universitat Duisburg-Essen
PA135 (#543)	Functional Capacity in Light Chain Amyloid Cardiomyopathy: Prognostic Value and Changes With Therapy	Olivier Clerc Mass General Brigham Inc
PA136 (#550)	Prognostic significance of circulating tumor cells assessed with next generation flow cytometry in patients with AL amyloidosis	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon
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molecular mechanisms characterizing the heart tissue of ALA and ATTR amyloidosis patients Early Development and Pre-Clinical Evaluation of a Fluorine-18 Labeled Peptide, p5+14, for the Detection of Amyloid Cardiomyopathy by PET/CT Imaging PB115 (#302) Uptake of Iodine (124) Evuzamitide in Patients with AL and ATTR Amyloidosis and Correlation with Echocardiographic Parameters PB116 (#362) Right heart dysfunction in patients with cardiac amyloidosis PB117 (#401) Comparative Analysis of Clinical and Echocardiographic Variations in Cardiac Amyloidosis Subtypes PB118 (#403) Prevalence of moderate-severe aortic stenosis in patients with wild-type transthyretin amyloidosis in a developing country. PB119 (#412) Clinical profile and outcome of cardiac amyloidosis in a middle-income country. PB120 (#416) Automatic quantification of AL and ATTR amyloidosis and individual sease burden using 1241-evuzamitide, a novel radiotracer PB121 (#418) Wild-type Transthyretin Cardiac Amyloidosis with Positive 18F-FDG/13N-ammonia Cardiac Positron Emission Tomography PB122 (#423) Relationship Between Myocardial Amyloid Load Measured by 1241-evuzamitide and Prognostic Staging Systems in Transthyretin Amyloid Cardiomyopathy PB123 (#424) Relationship Between Myocardial 1241-evuzamitide Uptake and Extracellular Volume Fraction: A Cardiac PET/MRI Study PB124 (#430) Longitudinal Contractile Diastasis: A Novel Myocardial Contraction-Relaxation Abnormality in Patients with Cardiac Amyloidosis on 1241-evuzamitide petr/MRI Study PB125 (#431) A reduced ejection fraction is associated with more severe myocardial Blood flow abnormality and biomarker elevation in ATTR cardiac amyloidosis PB126 (#434) Characterizing Renal Involvement in Light Chain Amyloidosis on 1241-evuzamitide PET/MRI Imaging PB126 (#434) Characterizing Renal Involvement in Light Chain Amyloidosis on 1241-evuzamitide PET/MRI Imaging	PB112	(#283)	Amyloidosis Evaluation among Patients with Aortic Stenosis Referred for Transcatheter Aortic Valve	
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	Molecular modeling of apoE interactions with amyloid-beta fibrils from human brain suggests a structural basis for apolipoprotein co-deposition with amyloids	Boston University Yohei Misumi
PC2 (#36)	Molecular modeling of apoE interactions with amyloid-beta fibrils from human brain suggests a structural basis for apolipoprotein co-deposition with amyloids Study of the binding of serum proteins to amyloid fibrils Analysis of SAA fragments digested by a lysosomal	Poston University Yohei Misumi Kumamoto Daigaku Kouhei Kawaguchi
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PC22 (#427)	Dual Amyloidosis Found on Transverse Carpel Ligament Biopsy: A Case Report	Ellen Lewis Alberta Government
PC23 (#447)	Diagnostic error and the importance of cardiac biopsy in the differentiation of amyloid protein: An atypical case of APOA1 amyloidosis	Diane Xavier De Ávila Fluminense Federal University
PC24 (#452)	Clinicopathologic and Outcome Characteristics of Renal Apolipoprotein C-II Amyloidosis	Samih Nasr Mayo Clinic
PC25 (#500)	Cathepsin K and Cystatin C are Co-deposited in Amyloids of Other Types	Jason Theis Mayo Clinic
PC26 (#510)	The Pavia Amyloidosis Research and Treatment Center's 25-year experience in molecular diagnostics for hereditary amyloidoses	Mario Nuvolone Universita degli Studi di Pavia

PC27 (#558)	Nodular pulmonary amyloidosis as a trigger for the diagnosis of sjogren's syndrome	Pedro Garibaldi Universidade de Sao Paulo
	PATIENT REPORTED OUTCOMES/QUALITY OF LIFE	
PC28 (#144)	RALTAN Study: Risk Assessment for acquired transthyretin Amyloid Neuropathyin Domino Liver Transplantation recipients	Ilias Kounis Universite Paris-Saclay
PC29 (#1)	Daily life in patients with hereditary transthyretin amyloidosis: a qualitative study.	Aina Isabel Gayá Barroso Health Research Institute of the Balearic Islands (IdISBa)
PC30 (#4)	Burden of transthyretin amyloid cardiomyopathy in treatment-naïve patients by heart failure severity: results from a large, non-interventional, real-world study	Francesco Cappelli Careggi University Hospital, Florence, Italy
PC31 (#5)	Predictors of disease burden in patients with untreated transthyretin amyloid cardiomyopathy: post hoc analysis of an international survey study	Francesco Cappelli Careggi University Hospital, Florence, Italy
PC32 (#6)	Caregiver burden by severity of patient's heart failure due to transthyretin amyloid cardiomyopathy: results from a large, non-interventional, real-world study	Francesco Cappelli Careggi University Hospital, Florence, Italy
PC33 (#7)	Predictors of burden in caregivers to patients with untreated transthyretin amyloid cardiomyopathy	Lucia Ponti University of Urbino Carlo Bo, Urbino, Italy
PC34 (#89)	Clinicopathological analyses of ATTRv amyloidosis after liver transplantation with cardiogenic cerebral embolism and subarachnoid hemorrhage	Shiori Yamakawa Kumamoto Daigaku
PC35 (#101)	Exercise Rehabilitation in Patients with Cardiac Amyloidosis, a Case study	Alexandros Briasoulis Ethniko kai Kapodistriako Panepistemio Athenon
PC36 (#106)	Health-related quality of life in hereditary transthyretin amyloidosis, a cross-sectional study	Delfina Cirelli Hospital Italiano de Buenos Aires
PC37 (#165)	Health-related quality-of-life, diagnosis and treatment experiences of AL amyloidosis patients	Solène Clavreul Myeloma Patients Europe
PC38 (#214)	Disease Burden of ATTR Amyloidosis Based on the SF- 36√2® Health Survey	Vaishali Sanchorawala Boston Medical Center
PC39 (#251)	A Platform for Federated Acquisition of Health-related Quality of Life Data in Systemic Amyloidosis	Tobias Dittrich Universitat Heidelberg
PC40 (#265)	The prevalence of frailty and association with disease severity in transthyretin amyloid cardiomyopathy	Elyn Montgomery Sisters of Charity of Australia
PC41 (#284)	Health-related Quality of Life as a predictor of Heart Failure Admission and Death in Patients with Amyloidosis	Konstantinos Sideris Utah System of Higher Education
PC42 (#316)	Quality of Life of Patients with Variant and Wild-Type Transthyretin Amyloidosis	Sabrina Rebello Amyloidosis Research Consortium

PC43 (#346)	Health-Related Quality of Life (HRQoL) in Patients With Mayo Stage IV Light Chain (AL) Amyloidosis Treated With Birtamimab Plus Standard of Care (SoC): Results From the VITAL Trial	Anita D'souza Medical College of Wisconsin
PC44 (#356)	Prevalence and outcome of coronary artery disease in cardiac amyloidosis: A retrospective Austrian multicenter study	Maria Ungericht Land Tirol
PC45 (#369)	Social Determinants of Health-Related Disparities in the Diagnosis of Transthyretin Cardiac Amyloidosis	Cindi Pankratova Pfizer Inc
PC46 (#375)	Experiences and benefits of setting up an Amyloidosis patient panel within an early phase clinical trials unit	James Rickard Richmond Research Institute
PC47 (#507)	Psychometric Validation of the Transthyretin Amyloidosis – Quality of Life (ATTR-QOL) Questionnaire	Kaitlin Lagasse UnitedHealth Group Inc
PC48 (#509)	Best Practices and Key Barriers for Amyloidosis Patient Care – a Representative Analysis of Multidisciplinary Amyloidosis Specialists, Referring Physicians, Patients, and Patient Advocacy Groups at US Specialized Amyloidosis Centers	Jose Nativi Mayo Clinic
PC49 (#521)	Bortezomib Induced Ocular Toxicities in AL Amyloidosis	Shahrier Hossain Dana-Farber Cancer Institute
	DDOCNOSIS ATTD	
PC50 (#25)	PROGNOSIS ATTR Fifty years of experience with hereditary transthyretin amyloidosis – first report from the SveATTR registry	Jonas Wixner Umea Universitet
PC51 (#45)	Performance of prognostic staging systems for Transthyretin Cardiac Amyloidosis in the disease modifying era.	Mat Maurer Columbia University
PC52 (#71)	Role of age and QRS duration as mortality predictors in a cohort of patients with transthyretin cardiac amyloidosis (ATTR-CA)	Cristhian Vicente Espinoza Romero Govemo do Estado de Sao Paulo
PC53 (#86)	Autonomic Dysfunction Severity and Cardiovascular Symptoms in ATTR Cardiac Amyloidosis: Insights from the COMPASS-31 Questionnaire	Ariel Weinsaft Columbia University
PC54 (#132)	Heart failure with preserved, mildly reduced, and reduced left ventricular ejection fraction in patients with transthyretin amyloid cardiomyopathy	Georgina Del Cisne Jadán Luzuriaga
PC55 (#137)	Clinical utility of the 6-minute walk distance in cardiac transthyretin amyloidosis	Vladimir Cejka Julius-Maximilians-Universitat Wurzburg
PC56 (#204)	Prospective, Real-World Data on the Characteristics, Treatment Patterns, and Outcomes of Patients With Transthyretin Amyloidosis: Design of the MaesTTRo Study	Julian D. Gillmore University of London

PC57 (#216)	Baseline predictors of major adverse events for transthyretin amyloidosis cardiomyopathy patients treated and untreated with disease-modifying therapy	Karan Shahi University of Calgary
PC58 (#236)	Predictors of Developing Renal Dysfunction Following Diagnosis of Transthyretin Cardiac Amyloidosis	Malcolm Mcdonald Cleveland Clinic
PC59 (#237)	Clinical Relevance of NT-Pro BNP at the Time of ATTR Cardiac Amyloidosis Diagnosis	Malcolm Mcdonald Cleveland Clinic
PC60 (#247)	Gamma-glutamyltransferase independently predicts all- cause mortality in transthyretin cardiac amyloidosis	Alberto Aimo Fondazione Toscana Gabriele Monasterio per la Ricerca Medica e di Sanita Pubblica
PC61 (#310)	The effect of tafamidis treatment on vascular aging in patients with ATTR cardiomyopathy. An observational study	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon
PC62 (#323)	The Impact of Frailty and Cardiac Amyloid Burden on 6-minute walk distance in ATTR cardiomyopathy.	Samsam Dahir Mass General Brigham Inc
PC63 (#351)	Transthyretin derived amyloid deposits in the atrium and the aortic valve: insights from multimodality evaluations and mid-term follow up	Atsushi Okada Kokuritsu Junkankibyo Kenkyu Center
PC64 (#353)	Transthyretin amyloidosis in Greece: Outcomes and changing diagnostic patterns	Efstathios Kastritis Ethniko kai Kapodistriako Panepistemio Athenon
PC65 (#373)	A Natural History Study of Transthyretin (ATTR) Amyloidosis: Trends in Overall Survival at a U.S. Referral Center	Andrew Staron Boston Medical Center
PC66 (#381)	Left Ventricular Obstruction in Transthyretin amyloid cardiomyopathy	Nerea Mora Ayestarán Universidad Autonoma de Madrid
PC67 (#389)	Impact of upgrading to cardiac resynchronization therapy or left bundle branch pacing in patients with ATTR cardiomyopathy	Belen Peiro-Aventin Universidad Autonoma de Madrid
PC68 (#395)	Natriuretic peptides, atrial fibrillation, and survival in patients with transthyretin cardiomyopathy receiving tafamidis treatment.	Dimitrios Bampatsias Columbia University
PC69 (#398)	Incidence of pacemaker implantation and prognostic value on all-cause mortality in wild-type transthyretin amyloidosis	Jens Skov Region Midtjylland
PC70 (#400)	Incidence and prognostic implications of atrial fibrillation in wild-type transthyretin amyloidosis cardiomyopathy	Jens Skov Region Midtjylland
PC71 (#428)	Evolution of Transthyretin Amyloid Cardiomyopathy Characteristics Over a 20 Year Period: A Multicenter Study of 1,168 Patients	Ahmad Masri Oregon Health & Science University

PC72 (#429)	A Multicenter Study of Real-World Outcomes of Tafamidis in Transthyretin Amyloid Cardiomyopathy	Ahmad Masri Oregon Health & Science University
PC73 (#450)	Predictors of early mortality in patients with wild-type transthyretin cardiac amyloidosis	Paolo Milani Universita degli Studi di Pavia
PC74 (#453)	Reduction in All-Cause Mortality in ATTR-CM Over a 20 year Period	Brett Sperry Saint Luke's Health System
PC75 (#478)	Transthyretin Amyloidosis: Does more than one pathogenic variant confer higher severity?	Larissa Bruscky Governo do Estado de Sao Paulo
PC76 (#487)	Can patisiran stop the progression of cardiac amyloidosis in patients with ATTRE109Q amyloidosis? Description of a monocentric experience.	Massimo Russo Universita degli Studi di Messina
PC77 (#493)	Role of Blood Biomarkers in Hereditary transthyretin amyloidosis: A Real-World Experience from a Single Center.	Massimo Russo Universita degli Studi di Messina
PC78 (#504)	Interventricular septum diameter as an echocardiographic predictor of thromboembolic events (TEEs) in patients with cardiac amyloidosis (CA) without atrial fibrillation diagnosis	Cristhian Vicente Espinoza Romero Governo do Estado de Sao Paulo
PC79 (#526)	Predictors of atrial fibrillation in Brazilian patients with transthyretin amyloidosis cardiomyopathy: Insights from the REACT registry.	Cristhian Vicente Espinoza Romero Governo do Estado de Sao Paulo
PC80 (#562)	Prevalence of frailty in a contemporary population of ATTR cardiac amyloidosis	Abigail Aronson Harvard University
PC81 (#566)	Survival in Wild-Type and Variant Transthyretin Amyloid Cardiomyopathy Over a 20-Year Period	Mazen Hanna Cleveland Clinic
PC82 (#569)	Lack of an association between baseline or follow-up transthyretin levels and all-cause mortality in ATTR cardiomyopathy patients on tafamidis	Abdullah Zoheb Azhar UPMC

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PC83 (#102)	Localized light chain (AL) amyloidosis presentation, treatment, and outcomes; an observational study.	Nirija Ranjit Anderson Queensland Government
PC84 (#139)	Hereditary AA Amyloidosis Caused by a Mutation in the SAA1 Promoter. Therapeutic Follow-up.	Jakub Sikora Univerzita Karlova
PC85 (#286)	Frequency and prognostic implications of translocation (11;14) in monoclonal immunoglobulin deposition disease	Samuel Rubinstein The University of North Carolina System
PC86 (#320)	Characteristics of Leukocyte Chemotactic Factor 2 (ALect2) Amyloidosis Patients from the Amyloidosis Research Consortium's 2023 Multi-Country Amyloidosis Community Survey	Sabrina Rebello Amyloidosis Research Consortium

PC87 (#386)	Risk factors associated with adverse renal outcomes in	Maria Vargas
	renal leukocyte chemotactic factor 2-associated amyloidosis (ALECT2)	Clinic
PC88 (#471)	Unveiling Natural history of Amyloidosis in Tuberculosis: A Novel Insight Using a Non-Genetic Mice Model	Akansha Garg Indian Institute of Technology Kanpur
PC89 (#556)	Clinicopathologic and Proteomic Features of Breast Amyloidosis	Michael Keeney Mayo Clinic
	TREATMENTS OF ATTR	
PC90 (#2)	Tafamidis 80 mg is more likely to improve disease measures than placebo in patients with transthyretin amyloid cardiomyopathy	Mazen Hanna Cleveland Clinic
PC91 (#9)	Tafamidis efficacy among octogenarian patients in the phase 3 ATTR-ACT and ongoing long-term extension study	Pablo Garcia-Pavia Universidad Autonoma de Madrid
PC92 (#13)	Safety and efficacy of tafamidis in Chinese patients with transthyretin amyloid cardiomyopathy	Zhuang Tian Peking Union Medical College Hospital, Beijing, China
PC93 (#14)	Real-World Effectiveness of High-Dose Tafamidis on Neurologic Disease Progression in Mixed-Phenotype Transthyretin Amyloid Cardiomyopathy	Nicholas S Streicher Society of Jesus
PC94 (#15)	Real-World Effectiveness of High-Dose Tafamidis on Neurologic Disease Progression in Mixed-Phenotype Variant Transthyretin Amyloid Cardiomyopathy	Nicholas S Streicher Society of Jesus
PC95 (#17)	Long-term tafamidis efficacy in patients with transthyretin amyloid cardiomyopathy by baseline left ventricular ejection fraction	Brian Drachman University of Pennsylvania Health System, Philadelphia, PA, USA
PC96 (#41)	SGLT-2- inhibitors in patients with cardiac wildtype ATTR-amyloidosis	Gina Barzen Freie Universitat Berlin
PC97 (#42)	Comparison of Combination Therapy (Tafamidis Plus Silencer) vs. Tafamidis Monotherapy in Transthyretin Amyloid Cardiomyopathy: Single Center Retrospective Cohort Study	Abdirahman Wardhere Columbia University
PC98 (#77)	Genotype vs phenotype and access to treatment in patients with hereditary transthyretin amyloidosis from a reference center in Brazil	Fabio De Souza Universidade Federal do Estado do Rio de Janeiro
PC99 (#82)	Treating Transthyretin Amyloidosis with CRISPR/Cas9 via LNP Delivery for in vivo TTR Knockdown	Emma Wang YolTech Therapeutics
PC100 (#116)	Open-label treatment data from the Swedish SveaTTR registry support the efficacy of diflunisal for hereditary transthyretin amyloidosis.	Jonas Wixner Umea Universitet
PC101 (#118)	Single Center Study of Long-Term Outcomes in ATTR-CM Patients treated with Tafamidis vs Acoramidis: Early findings at the dawn of a new era.	Dimitrios Bampatsias Columbia University

PC102 (#133)	Association between Patient Characteristics and Serum Neurofilament Light Chain Level in Patients with Transthyretin-Related Amyloid Cardiomyopathy	Bryton Davis Oregon Health & Science University
PC103 (#151)	Enabling the development of serum TTR as a biomarker for treatment of ATTR amyloidosis	Julian D. Gillmore University of London
PC104 (#152)	Design and rationale for MAGNITUDE, a phase 3, randomized, placebo-controlled trial of NTLA-2001, a CRISPR-based gene editing therapy, in patients with transthyretin amyloidosis with cardiomyopathy (ATTR-CM)	Mathew S. Maurer Columbia University
PC105 (#171)	Effect and Safety of Patisiran in Patients with Hereditary Transthyretin Amyloidosis with Polyneuropathy and Chronic Kidney Disease	Julien Dang CHU Bicêtre, Université Paris- Saclay, France
PC106 (#174)	Neuropathy Impairment and Nutritional Status with Eplontersen in Patients with Hereditary Transthyretin- Mediated Amyloidosis	Jonas Wixner Umea Universitet
PC107 (#10)	Survival in a real-world cohort of patients with mixed phenotype transthyretin amyloidosis treated with tafamidis: an analysis from the Transthyretin Amyloidosis Outcomes Survey	Jonas Wixner
PC108 (#176)	History of Polyneuropathy and Musculoskeletal Manifestations in Patients with Transthyretin-Mediated Amyloidosis with Cardiomyopathy in APOLLO-B	Martha Grogan Mayo Clinic
PC109 (#182)	A unified analysis of the impact of rapid TTR knockdown by RNAi therapeutics across transthyretin-mediated (ATTR) amyloidosis studies	Martha Grogan MayoClinic
PC110 (#190)	Rationale and Design of ACT-EARLY, the Acoramidis Transthyretin Amyloidosis Prevention Trial	Pablo Garcia-Pavia Universidad Autonoma de Madrid
PC111 (#196)	Arginine, as a possible amino acid for prevention of ATTR amyloidosis	Atsushi Fukunari Gakko Hojin Kyushu Bunka Gakuen
PC112 (#202)	Eplontersen for Hereditary Transthyretin Amyloidosis With Polyneuropathy: An Exploratory Analysis of Treatment Effect in Male and Female Patients	Márcia Waddington Cruz CEPARM, University Hospital, Federal University of Rio de Janeiro
PC113 (#217)	Magnitude and timing of change in monitoring parameters for transthyretin amyloidosis cardiomyopathy patients treated and untreated with tafamidis	Karan Shahi University of Calgary
PC114 (#218)	Transthyretin amyloidosis cardiomyopathy disease staging systems for patients treated with tafamidis therapy	Karan Shahi University of Calgary
PC115 (#219)	Determinants guiding clinical decision-making for treatment initiation in transthyretin amyloidosis cardiomyopathy patients in canada	Karan Shahi University of Calgary
PC116 (#227)	ALXN2220: targeted immune activation in patients undergoing antibody-mediated cardiac ATTR amyloid depletion	Peter Christian Kahr Neurimmune

PC117 (#228)	Reduction of amyloid deposition by patisiran —Evaluation by gastroduodenal biopsy and amyloid 11C-Pittsburgh compound B PET	Shinji Masuko Shinshu Daigaku
PC118 (#230)	Eplontersen for Hereditary Transthyretin Amyloidosis With Polyneuropathy: An Exploratory Analysis in Patients With the V30M TTR Variant and Early-Onset or Late-Onset Disease	Julian D. Gillmore University of London
PC119 (#233)	Single-center one-year outcome of Vutrisiran treatment in hereditary transthyretin amyloidosis	Helena Pemice Freie Universitat Berlin
PC120 (#241)	Racial/Ethnic Differences in Treatment Utilization for ATTR Amyloidosis	Alexandra Greatsinger Analysis Group Inc
PC121 (#245)	Safety and efficacy of levosimendan in patients with cardiac amyloidosis	Alberto Aimo Fondazione Toscana Gabriele Monasterio per la Ricerca Medica e di Sanita Pubblica
PC122 (#279)	Higher Risk of Mortality in Previously Hospitalized Patients: Insights from ATTRibute-CM	Ahmad Masri Oregon Health & Science University
PC123 (#291)	Prediction of cardiac ATTR depletion by NI006 using mechanistic PK/PD modeling	Aubin Michalon Neurimmune AG
PC124 (#292)	Treatment-related Early Increase in Serum TTR is Associated with Lower Cardiovascular Mortality in ATTR- CM: Insights from ATTRibute-CM	Amrut Ambardekar University of Colorado System
PC125 (#295)	Improved Health-Related Quality of Life in Acoramidis- Treated Patients with ATTR-CM, Demonstrated by Improvements in KCCQ Scores	Marianna Fontana University of London
PC126 (#300)	ATTRibute-CM: ITT Sensitivity Analysis and Sub-Analysis Comparing Acoramidis and Placebo in Stage 4 CKD	Steen Hvitfeldt Poulsen Region Midtjylland
PC127 (#303)	Acoramidis significantly improves NT-proBNP indices that indicate ATTR-CM disease progression and predict subsequent mortality: Insights from the ATTRibute-CM Study	Pablo Garcia-Pavia Universidad Autonoma de Madrid
PC128 (#315)	Acoramidis May Improve Cardiac Function and Promote Regression in ATTR-CM: Data From the ATTRibute-CM Cardiac Magnetic Resonance (CMR) Substudy	Yousuf Razvi University of London
PC129 (#317)	Safety of Direct Current Cardioversion Without Routine Transesophageal Echocardiography in Patients with Cardiac Amyloidosis	Osnat Itzhaki Ben Zadok Mass General Brigham Inc
PC130 (#321)	Early Increase in Serum Transthyretin Level is an Independent Predictor of Improved Survival in ATTR Cardiomyopathy: Insights From Acoramidis Phase 3 Study ATTRibute-CM	Mat Maurer Columbia University
PC131 (#344)	Arrhythmia burden and progression in ATTR-CM patients treated and untreated with tafamidis	Mannat Dhillon University of Calgary

PC132 (#349)	Acoramidis Improves Clinical Outcomes in Transthyretin Amyloid Cardiomyopathy	Daniel Judge Medical University of South Carolina
PC133 (#374)	Disease progression in patients with wild-type transthyretin cardiomyopathy on treatment with tafamidis: prevalence and clinical correlates.	Giulio Sinigiani Universita degli Studi di Padova
PC134 (#396)	ATTR V30M Nephropathy: Implications of switching anti- amyloid agents	Joana Tavares Governo da Republica Portuguesa
PC135 (#399)	Early tolerance of SGLT2i therapy in patients with ATTR V30M amyloidosis: staging and scoring for a standard of care evaluation across different clinical phenotypes	João Bessa E Silva Governo da Republica Portuguesa
PC136 (#402)	Health-Related Quality of Life in Patients with Symptomatic Transthyretin Amyloid Cardiomyopathy Treated with Acoramidis: An EQ-5D Analysis From the ATTRibute-CM Study	Mazen Hanna Cleveland Clinic
PC137 (#409)	SGLT2 Inhibitors for Transthyretin Amyloid Cardiomyopathy: A Propensity Score-Matched Analysis	Frederick M. Lang Columbia University
PC138 (#414)	The Amyloidosis Forum: Five Years of Progress for the Public-Private Partnership to Advance Drug Development for Amyloidosis	Kristen Hsu Amyloidosis Research Consortium
PC139 (#441)	A phase III randomized study of doxycycline and tauroursodeoxycholic acid (Doxy/TUDCA) plus standard supportive therapy versus standard supportive therapy alone in cardiac amyloidosis caused by transthyretin	Paolo Milani Universita degli Studi di Pavia
PC140 (#465)	Regression of amyloid load in subcutaneous fat tissue of hereditary transthyretin amyloidosis patients during treatment with patisiran	Hendrea Tingen Rijksuniversiteit Groningen
PC141 (#498)	Establishing amyloidosis clinics at a university hospital in the Northwest of Germany	Christoph Mronga Stadt Oldenburg
PC142 (#518)	Descriptive characteristics of patients diagnosed with transthyretin amyloidosis in the commercial and Medicare populations	Neela Kumar AstraZeneca PLC
PC143 (#519)	Clinical Trial Design of AT-02 Phase 2 Open Label Extension Study in Systemic Amyloidosis	Spencer Guthrie Attralus
PC144 (#538)	Predictors of early mortality after initiation of tafamidis therapy.	Prem Soman University of Pittsburgh
PC145 (#540)	Comparison of treatment responders versus non- responders in transthyretin amyloid cardiomyopathy	Joban Vaishnav Johns Hopkins University
	TREATMENTS OF OTHER MORE RARE AMYLOIDOSES	
PC146 (#199)	Potential use of Multifunctional Therapeutic Cyclodextrin- Appended Dendrimer Complex for Systemic & Localized Amyloidosis Treatment	Hirofumi Jono Kumamoto Daigaku
PC147 (#285)	Outcomes of non-AL monoclonal gammopathies of renal significance with clone-directed therapy	Samuel Rubinstein
	Significance with cione-unected therapy	The University of North Carolina System
PC148 (#371)	Clinical presentation and treatment of genitourinary amyloidosis: A single referral center 30-year experience	Lisa Mendelson Boston University