

Concepts for the Development of Person-Centered, Digitally Enabled, Artificial Intelligence—Assisted ARIA Care Pathways (ARIA 2024)



Jean Bousquet, MD^{a,b,c}, Holger J. Schünemann, MD^{b,d,e}, Bernardo Sousa-Pinto, MD^{f,g}, Torsten Zuberbier, MD^{a,b,d}, Alkis Togias, MD^h, Boleslaw Samolinski, MDⁱ, Anna Bedbrook, BSc^c, Wienczyslaw Czarlewski, MD^{c,j}, Martin Hofmann-Apitius, PhD^k, Justyna Litynska, MSc^{l,m}, Rafael José Vieira, MD^{f,g}, Josep M. Anto, MD^{n,o,p}, Joao A. Fonseca, MD^{f,g}, Jan Brozek, MD^d, Antonio Bognanni, MD^q, Luisa Brussino, PhD^{r,s}, G. Walter Canonica, MD^{t,u}, Ivan Cherrez-Ojeda, MD^{v,w}, Alvaro A. Cruz, MD^x, Leticia de las Vecillas, MD^y, Mark Dykewicz, MD^z, Bilun Gemicioglu, MD^{aa}, Mattia Giovannini, MD^{bb,cc}, Tari Haahtela, MD^{dd}, Marc Jacobs, PhD^k, Cristina Jacomelli, MSc^{ee}, Ludger Klimek, MD^{ff,gg}, Violeta Kvedariene, MD^{hh,ii}, Desiree E. Larenas-Linnemann, MD^{jj}, Gilles Louis, PhD^{kk}, Olga Lourenço, PhD^{ll}, Lucas Leemann, PhD^{mm}, Mario Morais-Almeida, MDⁿⁿ, Ana Luisa Neves, MD^{oo}, Kari C. Nadeau, MD^{pp}, Artur Nowak, MSc^{l,m}, Yuliia Palamarchuk, PhD^{qq}, Susanna Palkonen, PhD^{rr}, Nikolaos G. Papadopoulos, MD^{ss}, Elena Parmelli, PhD^{t,u}, Ana Margarida Pereira, MD^{f,tt,uu}, Oliver Pfaar, MD^{vv}, Frederico S. Regateiro, MD^{ww,xx,yy,bbb}, Marine Savouré, PhD^{zz}, Luis Taborda-Barata, MD^{aaa,bbb}, Sanna K. Toppila-Salmi, MD^{ccc,ddd}, Maria J. Torres, MD^{eee}, Arunas Valiulis, MD^{fff,ggg}, Maria Teresa Ventura, MD^{hhh,iii}, Sian Williams, PhD^{jjj}, Juan J. Yepes-Núñez, MD^{kkk,lll}, Arzu Yorgancioglu, MD^{mmm}, Luo Zhang, MDⁿⁿⁿ, Jaron Zuberbier, MD^{ooo}, Amir Hamzah Abdul Latiff, MD^{ppp}, Baharudin Abdullah, MD^{qqq}, Ioana Agache, MD^{rrr}, Mona Al-Ahmad, MD^{sss}, Maryam Ali Al-Nesf, MD^{ttt}, Nada A. Al Shaikh, MD^{uuu}, Rita Amaral, PhD^{f,g,vvv,www}, Ignacio J. Ansotegui, MD^{xxx}, Julijana Asllani, MD^{yyy}, Maria Cristina Balotro-Torres, MD^{zzz}, Karl-Christian Bergmann, MD^{a,b}, Jonathan A. Bernstein, MD^{aaaa}, Carsten Bindselev-Jensen, MD^{bbbb}, Michael S. Blaiss, MD^{cccc}, Cristina Bonaglia, PhD^{dddd}, Matteo Bonini, PhD^{eeee,ffff,gggg}, Isabelle Bossé, PhD^{hhhh}, Fulvio Braido, PhD^{iiii,jjjj}, Fernan Caballero-Fonseca, MD^{kkkk}, Paulo Camargos, MD^{llll}, Pedro Carreiro-Martins, MD^{mmmm,nnnn}, Thomas Casale, MD^{oooo}, José-Antonio Castillo-Vizueté, MD^{pppp,qqqq}, Lorenzo Cecchi, MD^{rrr}, Maria do Ceu Teixeira, MD^{ssss,tttt}, Yoon-Seok Chang, MD^{uuuu}, Claudia Chaves Loureiro, MD^{vvvv,www}, George Christoff, PhD^{xxxx}, Giorgio Ciprandi, MD^{yyyy}, Ieva Cirule, MD^{zzzz}, Jaime Correia-de-Sousa, MD^{aaaaa}, Elisio M. Costa, PhD^{bbbbb}, Biljana Cvetkovski, PhD^{ccccc}, Govert de Vries, PhD^{ddddd}, Stefano Del Giacco, MD^{eeeee}, Philippe Devillier, MD^{fffff}, Dejan Dokic, MD^{ggggg}, Habib Douagui, MD^{hhhhh}, Stephen R. Durham, MDⁱⁱⁱⁱⁱⁱ

^aInstitute of Allergology, Charité—Universitätsmedizin Berlin, Corporate Member of Freie Universität Berlin and Humboldt-Universität zu Berlin, Berlin, Germany

^bFraunhofer Institute for Translational Medicine and Pharmacology ITMP, Immunology and Allergology, Berlin, Germany

^cARIA, Montpellier, France

^dDepartment of Health Research Methods, Evidence, and Impact & Department of Medicine, McMaster University, Hamilton, ON, Canada

^eDepartment of Biomedical Science, Humanitas University, Pieve Emanuele, Milan, Italy

^fMEDCIDS—Department of Community Medicine, Information and Health Decision Sciences, Faculty of Medicine, University of Porto, Porto, Portugal

^gCINTESIS@RISE—Health Research Network, Faculty of Medicine, University of Porto, Porto, Portugal

^hDivision of Allergy, Immunology, and Transplantation (DAIT), National Institute of Allergy and Infectious Diseases, NIH, Bethesda, Md

ⁱDepartment of Prevention of Environmental Hazards, Allergology and Immunology, Medical University of Warsaw, Warsaw, Poland

^jMedical Consulting Czarlewski, Levallois, France

^kDepartment of Bioinformatics, Fraunhofer Institute for Algorithms and Scientific Computing (SCAI), Bonn, Germany

^lEvidence Prime, Krakow, Poland

^mGynecological and Obstetric Polyclinic, District Hospital, Białystok, Poland

ⁿISGlobal, Barcelona Institute for Global Health, Barcelona, Spain

^oDepartment of Experimental Sciences, Universitat Pompeu Fabra (UPF), Barcelona, Spain

^pDepartment of Epidemiology & Public Health, CIBER Epidemiología y Salud Pública (CIBERESP), Barcelona, Spain

^qDepartment of Health Research Methods, Evidence, and Impact & Department of Medicine, Evidence in Allergy group, McMaster University, Hamilton, ON, Canada

^rDepartment of Medical Sciences, University of Torino, Torino, Italy

^sAllergy and Clinical Immunology Unit, Mauriziano Hospital, Torino, Italy

^tDepartment of Biomedical Sciences, Humanitas University, Pieve Emanuele, Milan, Italy

^uAsthma & Allergy Unit, IRCCS Humanitas Research Hospital, Rozzano, Milan, Italy

^vResearch Center 'Respiralab', Universidad Espíritu Santo, Samborondón, Ecuador

^wDepartment of Allergology & Pulmonology, Respiralab Research Group, Guayaquil, Guayas, Ecuador

^xFundação ProAR, Federal University of Bahia and GARD/WHO Planning Group, Salvador, Bahia, Brazil

^yDepartment of Allergy, Hospital La Paz Institute for Health Research (IdiPAZ), Madrid, Spain

^zSection of Allergy and Immunology, Saint Louis University School of Medicine, Saint Louis, Mo

^{aa}Department of Pulmonary Diseases, and Institute of Pulmonology & Tuberculosis, Istanbul University-Cerrahpaşa, Cerrahpaşa Faculty of Medicine, Istanbul, Turkey

^{bb}Department of Health Sciences, University of Florence, Florence, Italy

^{cc}Allergy Unit, Meyer Children's Hospital IRCCS, Florence, Italy

Maria Lourdes Enecilla, MD^{jjjjj}, Alessandro Fiocchi, MD^{kkkkk}, Wytse J. Fokkens, MD^{lllll}, Jean-François Fontaine, MD^{mmmmm}, Radoslaw Gawlik, MDⁿⁿⁿⁿⁿ, Jose E. Gereda, MD^{ooooo}, Sara Gil-Mata, MD^{f.g}, Antonio F.M. Giuliano, PhD^{ppppp}, Maia Gotua, MD^{qqqqq}, Brigita Gradauskiene, MD^{rrrrr}, Maria Antonieta Guzman, MD^{sssss}, Elham Hossny, MD^{ttttt}, Martin Hrubisko, MD^{uuuuu}, Tomohisa Iinuma, MD^{vvvvv}, Carla Irani, MD^{wwwww}, Zhanat Ispayeva, MD^{xxxxx}, Juan Carlos Ivancevich, MD^{yyyyy}, Tuomas Jartti, MD^{zzzzz,aaaaa}, Miloš Jeseňák, MD^{bbbbbb,ccccc}, Kaja Julge, MD^{ddddd}, Marek Jutel, MD^{eeeeee,ffffff}, Igor Kaidashev, MD^{gggggg}, Kazi S. Bennoor, MD^{hhhhh}, Nicolai Khaltaev, MDⁱⁱⁱⁱⁱⁱ, Bruce Kirenga, MD^{jjjjj}, Helga Kraxner, MD^{kkkkk}, Inger Kull, PhD^{lllll,mmmmm}, Marek Kulus, MDⁿⁿⁿⁿⁿ, Piotr Kuna, MD^{ooooo}, Maciej Kupczyk, MD^{ooooo}, Andriy Kurchenko, MD^{ppppp}, Stefania La Grutta, MD^{qqqqq}, Stephen Lane, MD^{rrrrr,sssss}, Neven Miculinic, MD^{ttttt}, Sang Min Lee, MD^{uuuuu}, Lan Le Thi Tuyet, MD^{vvvvv}, Battur Lkhagvaa, MD^{wwwww}, Renaud Louis, MD^{xxxxx,yyyyy}, Bassam Mahboub, MD^{zzzzz,aaaaa}, Mika Makela, MD^{dd}, Michael Makris, MD^{bbbbbb}, Marcus Maurer, MD^{a,b}, Eric Melén, MD^{lllll,mmmmm}, Branislava Milenkovic, MD^{ccccc}, Youssef Mohammad, MD^{ddddd,eeeeee}, Marcin Moniuszko, MD^{ffffff}, Stephen Montefort, MD^{gggggg}, Andre Moreira, MD^{hhhhh,iiiiii,jjjjj}, Pablo Moreno, MD^{kkkkk}, Joaquim Mullol, MD^{lllll,mmmmm}, Rachel Nadif, PhD^{nnnnn,ooooo}, Alla Nakonechna, MD^{ppppp,qqqqq}, Cecilia Gretchen Navarro-Locsin, MD^{rrrrr,sssss}, Hugo E. Neffen, MD^{ttttt}, Kristof Nekam, MD^{uuuuu}, Marek Niedoszytko, MD^{vvvvv}, Elizabete Nunes, MD^{wwwww}, Dieudonné Nyembue, MD^{xxxxx}, Robyn O'Hehir, MD^{yyyyy}, Markus Ollert, MD^{zzzzz,aaaaa}, Ken Ohta, MD^{bbbbbb}, Yoshitaka Okamoto, MD^{ccccc,ddddd}, Kimihiro Okubo, MD^{eeeeee}, Heidi Olze, MD^{ooo}, Mahesh Anand Padukudru, MD^{ffffff}, Oscar Palomares, PhD^{gggggg}, Isabella Pali-Schöll, MD^{hhhhh,iiiiii}, Petr Panzner, MD^{jjjjj}, Kati Palosuo, MD^{kkkkk}, Hae S. Park, MD^{lllll}, Giovanni Passalacqua, MD^{mmmmm}, Vincenzo Patella, MD^{nnnnn,ooooo,ppppp}, Ruby Pawankar, MD^{qqqqq}, Benoît Pétré, MD^{kk}, Constantinos Pitsios, MD^{rrrrr}, Davor Plavec, MD^{sssss,ttttt}, Todor A. Popov, MD^{uuuuu}, Francesca Puggioni, MD^{vvvvv}, Santiago Quirce, MD^y, Filip Raciborski, MDⁱ, Agn  Ramonait , MD^{wwwww,xxxxx}, Marysia Recto, MD^{yyyyy}, Susana Repka-Ramirez, MD^{zzzzz}, Graham Roberts, MD^{aaaaa,bbbbbb,ccccc}, Karla Robles-Velasco, MD^{v,w}, Nicolas Roche, MD^{ddddd,eeeeee,ffffff}, Monica Rodriguez-Gonzalez, MD^{ggggg}, Joel A. Romualdez, MD^{hhhhh}, Menachem Rottem, MD^{iiiiii,jjjjj}, Philip W. Rouadi, MD^{kkkkk,lllll}, Marianella Salapatras, MD^{mmmmm}, Joaquin Sastre, MDⁿⁿⁿⁿⁿ, Faradiba S. Serpa, MD^{ooooo}, Zineb Sayah, MD^{ppppp}, Nicola Scichilone, MD^{qqqqq}, Gianenrico Senna, MD^{rrrrr}, Juan Carlos Sisul, MD^{sssss}, Dirceu Sol , MD^{ttttt}, Manuel E. Soto-Martinez, MD^{uuuuu}, Milan Sova, MD^{vvvvv}, Olga Sozinova, PhD^{wwwww}, Katarina Stevanovic, PhD^{a,b}, Charlotte Suppli Ulrik, MD^{xxxxx,yyyyy}, Anna Szylling, MDⁱ, Frances M. Tan, MD^{zzzzz}, Pongsakorn Tantilipikorn, MD^{aaaaa}, Ana Todo-Bom, MD^{bbbbbb}, Vesna Tomic-Spiric, MD^{ccccc,ddddd}, Vladyslav Tsaryk, MD^{ppppp}, Ioanna Tsiligianni, PhD^{eeeeee,ffffff}, Marilyn Urrutia-Pereira, MD^{ggggg},

^{dd}Skin and Allergy Hospital, Helsinki University Hospital, and University of Helsinki, Helsinki, Finland

^{ee}“Respiriamo Insieme” Association, Asthma & Allergy Center, Padova, Italy

^{ff}Department of Otolaryngology, Head & Neck Surgery, Universit tsmedizin Mainz, Mainz, Germany

^{gg}Center for Rhinology and Allergology, Allergology & Rhinology Department, Wiesbaden, Germany

^{hh}Institute of Clinical Medicine, Clinic of Chest Diseases and Allergology, Faculty of Medicine, Vilnius University, Vilnius, Lithuania

ⁱⁱInstitute of Biomedical Sciences, Department of Pathology, Faculty of Medicine, Vilnius University, Vilnius, Lithuania

^{jj}Center of Excellence in Asthma and Allergy, M dica Sur Clinical Foundation and Hospital, M xico City, M xico

^{kk}Department of Public Health, University of Li ge, Li ge, Belgium

^{ll}Faculty of Health Sciences and CICS—UBI, Health Sciences Research Centre, University of Beira Interior, Covilh , Portugal

^{mm}Department of Political Science, University of Z rich, Z rich, Switzerland

ⁿⁿAllergy Center, CUF Descobertas Hospital, Lisbon, Portugal

^{oo}Global Digital Health Unit, Department of Primary Care and Public Health, Imperial College, London, United Kingdom

^{pp}Department of Environmental Health, Harvard TH Chan School of Public Health and Department of Medicine, Division of Allergy and Inflammation, Beth Israel Deaconess Hospital, Boston, Mass

^{qq}Finnish Meteorological Institute (FMI), Atmospheric Composition Research, Helsinki, Finland

^{rr}EFA European Federation of Allergy and Airways Diseases Patients' Associations, Brussels, Belgium

^{ss}Allergy Department, 2nd Pediatric Clinic, University of Athens, Athens, Greece

^{tt}PaCeIT—Patient Centered Innovation and Technologies, Center for Health Technology and Services Research (CINTESIS), Faculty of Medicine, University of Porto, Porto, Portugal

^{uu}Allergy Unit, Instituto and Hospital CUF, Porto, Portugal

^{vv}Section of Rhinology and Allergy, Department of Otorhinolaryngology, Head and Neck Surgery, University Hospital Marburg, Philipps-Universit t Marburg, Marburg, Germany

^{ww}Allergy and Clinical Immunology Unit, Centro Hospitalar e Universit rio de Coimbra, Coimbra, Portugal

^{xx}Center for Innovative Biomedicine and Biotechnology (CIBB), Faculty of Medicine, University of Coimbra, Coimbra, Portugal

^{yy}Institute of Immunology, Faculty of Medicine, University of Coimbra, Coimbra, Portugal

^{zz}Universit  de Rennes, Inserm, EHESP, Irset (Institut de recherche en sant , environnement et travail)—UMR_S 1085, Rennes, France

^{aaa}Department of Immunoallergology, Cova da Beira University Hospital Centre, Covilh , Portugal

^{bbb}UBIAir—Clinical & Experimental Lung Centre and CICS-UBI Health Sciences Research Centre, University of Beira Interior, Covilh , Portugal

^{ccc}Department of Allergy, Skin and Allergy Hospital, Inflammation Center, Helsinki University Hospital and University of Helsinki, Helsinki, Finland

Marylin Valentin Rostan, MD^{hhhhhhhhhh}, Mikhail Sofiev, PhD^{qq}, Erkka Valovirta, MDⁱⁱⁱⁱⁱⁱⁱ, Michiel Van Eerd, PhD^{ddddd}, Eric Van Ganse, MD^{jjjjjjj}, Tuula Vasankari, MD^{kkkkkkkkkk, llllllll}, Pakit Vichyanond, MD^{mmmmmmmmmm}, Giovanni Vieg, MDⁿⁿⁿⁿⁿⁿⁿⁿⁿⁿ, Dana Wallace, MD^{oooooooooo}, De Yun Wang, MD^{pppppppppp}, Susan Waserman, MSc^{qqqqqqqqqq}, Gary Wong, MD^{rrrrrrrr}, Margitta Worm, MD^{ssssssssss}, Osman M. Yusuf, MD^{ttttttttt}, Fares Zaitoun, MD^{uuuuuuuuuu}, and Mihaela Zidarn, MD^{vvvvvvvvvv, wwwwwwwwwwww}

Berlin, Bonn, Mainz, Wiesbaden, and Marburg, Germany; Montpellier, Levallois, Rennes, La Rochelle, Suresnes, Reims, Villejuif, Paris, and Lyon, France; Hamilton, ON, Canada; Torino, Milan, Florence, Padova, Bari, Rome, Genoa, Cagliari, Prato, Genova, Palermo, Salerno, Naples, Verona, Pisa, Italy; Brussels and Liège, Belgium; Braga, Coimbra, Covilhã, Lisbon, and Porto, Portugal; Bethesda, Md; Białystok, Białystok, Gdansk, Katowice, Krakow, Lodz, and Warsaw, Wrocław, Poland; Barcelona, Bilbao, Madrid, and Malaga, Spain; Guayas and Samborondón, Ecuador; Uruguiana, Bahia, Belo Horizonte, Espírito Santo, and São Paulo, Brazil; Saint Louis, Mo; Istanbul, Turkey; Helsinki, Kuopio, Oulu, and Turku, Finland; Kaunas, Klaipeda, and Vilnius, Lithuania; México City, Mexico; Zürich, Switzerland; Isle of Wight, Liverpool, London, Sheffield, Southampton, Aberdeen and Edinburgh, United Kingdom; Boston, Mass; Bogotá, DC, Colombia; Beijing and Hong Kong, SAR, China; Kuala Lumpur, and Kubang Kerian, Kelantan, Malaysia; Brasov, Romania; Kuwait City and Salmiya, Kuwait; Doha, Qatar; Dammam, Saudi Arabia; Stockholm and Uppsala, Sweden; Tirana, Albania; Mandaluyong, Manila, and Quezon City, Philippines; Cincinnati, Ohio; Copenhagen and Odense, Denmark; Augusta, Ga; Caracas, Venezuela; Fort Lauderdale and Tampa, Fla; Praia, Cabo Verde; Seoul, Suwon, and Cheonan, Republic of Korea; Sofia, Bulgaria; Riga, Latvia; Sydney, NSW, and Melbourne, VIC, Australia; Amsterdam and Geldermalsen, the Netherlands; Skopje, Republic of Macedonia; Algiers, Algeria; Lima, Peru; Tbilisi, Georgia; Santiago, Chile; Cairo, Egypt; Bratislava, Slovakia; Chiba and Tokyo, Japan; Beirut and Lebanon; Almaty, Kazakhstan; Buenos Aires and Santa Fe, Argentina; Tartu, Estonia; Kyiv and Poltava, Ukraine; Dhaka, Bangladesh; Geneva, Switzerland; Kampala, Uganda; Budapest, Hungary; Dublin, Ireland; Zagreb and Osijek, Croatia; Hô-Chi-Minh City, Vietnam; Ulan Bator, Mongolia; Dubai and Sharjah, UAE; Athens and Heraklion, Greece; Belgrade, Serbia; Damascus and Latakia, Syria; Msida MSD, Malta; Maputo, Mozambique; Kinshasa, Democratic Republic of the Congo; Esch-sur-Alzette, Luxembourg; Karnataka, India; Vienna, Austria; Prague and Brno, Czech Republic; Nicosia, Cyprus; Asuncion and San Lorenzo, Paraguay; Afulaa nnd Haifa, Israel; Rabat, Morocco; San Jose, Costa Rica; Bangkok, Thailand; Belgrade, Serbia; Montevideo, Uruguay; Singapore, Republic of Singapore; Islamabad, Pakistan; and Ljubljana, Slovenia

The traditional healthcare model is focused on diseases (medicine and natural science) and does not acknowledge patients' resources and abilities to be experts in their own lives based on their lived experiences. Improving healthcare safety, quality, and coordination, as well as quality of life, is an important aim in the care of patients with chronic conditions. Person-centered care needs to ensure that people's values and

preferences guide clinical decisions. This paper reviews current knowledge to develop (1) digital care pathways for rhinitis and asthma multimorbidity and (2) digitally enabled, person-centered care.¹ It combines all relevant research evidence, including the so-called real-world evidence, with the ultimate goal to develop digitally enabled, patient-centered care. The paper includes (1) Allergic Rhinitis and its Impact on Asthma

^{ddd}Department of Otorhinolaryngology, University of Eastern Finland and the North Savo Wellbeing Services County, Kuopio, Finland

^{eee}Allergy Unit, Málaga Regional University Hospital of Málaga, Malaga University, ARADyAL, Malaga, Spain

^{fff}Clinic of Children's Diseases, Institute of Clinical Medicine, Medical Faculty of Vilnius University, Vilnius, Lithuania

^{ggg}Clinic of Asthma, Allergy, and Chronic Lung Diseases, Asthma & Allergy Department, Vilnius, Lithuania

^{hhh}Allergy and Clinical Immunology, University of Bari Medical School, Bari, Italy

ⁱⁱⁱInstitute of Sciences of Food Production, National Research Council (ISPA-CNR), Bari, Italy

^{jjj}International Primary Care Respiratory Group IPCRG, Edinburgh, United Kingdom

^{kkk}School of Medicine, Universidad de los Andes, Bogotá, DC, Colombia

^{lll}Pulmonology Service, Internal Medicine Section, Fundación Santa Fe de Bogotá, University Hospital, Bogotá, DC, Colombia

^{mmm}Department of Pulmonary Diseases, Celal Bayar University, Faculty of Medicine, Manisa, Turkey

ⁿⁿⁿDepartment of Otolaryngology, Head and Neck Surgery, Beijing TongRen Hospital and Beijing Institute of Otolaryngology, Beijing, China

^{ooo}Department of Otorhinolaryngology, Charité—Universitätsmedizin Berlin, Berlin, Germany

^{ppp}Allergy & Immunology Centre, Pantai Hospital Kuala Lumpur, Kuala Lumpur, Malaysia

^{qqq}Department of Otorhinolaryngology—Head and Neck Surgery, School of Medical Sciences, Universiti Sains Malaysia, Kubang Kerian, Kelantan, Malaysia

^{rrr}Faculty of Medicine, Transylvania University, Brasov, Romania

^{sss}Microbiology Department, College of Medicine, Kuwait University, Kuwait City, Kuwait

^{ttt}Adult Allergy and Immunology Division—Hamad Medical Corporation, Doha, Qatar

^{uuu}Otolaryngology Head & Neck Surgery Department, Mouwasat Hospital, Dammam, Saudi Arabia

^{vvv}Department of Cardiovascular and Respiratory Sciences, Porto Health School, Polytechnic Institute of Porto, Porto, Portugal

^{www}Department of Women's and Children's Health, Paediatric Research, Uppsala University, Uppsala, Sweden

^{xxx}Department of Allergy and Immunology, Hospital Quironsalud Bizkaia, Bilbao, Spain

^{yyy}Department of Internal Medicine, University of Medicine, Tirana, Albania

^{zzz}Section of Allergy, Asthma and Immunology, University of Santo Tomas Hospital, Manila, Philippines

^{aaa}Division of Immunology, Allergy and Rheumatology, Department of Medicine, University of Cincinnati College of Medicine, Cincinnati, Ohio

^{bbb}Department of Dermatology and Allergy Center, Odense Research Centre for Anaphylaxis (ORCA), Odense University Hospital, Odense, Denmark

^{ccc}Department of Pediatrics, Medical College of Georgia at Augusta University, Augusta, Ga

^{ddd}Institute "Bona Sforza", University for Linguistic Mediators, Bari, Italy

Abbreviations used

AI- Artificial intelligence
AIRWAYS-ICPs- Integrated Care Pathways for Airway Diseases
AIT- Allergen immunotherapy
AR- Allergic rhinitis
ARIA- Allergic Rhinitis and its Impact on Asthma
CARAT- Control of Allergic Rhinitis and Asthma Test
CATALYSE- Climate Action To Advance HeaLthY Societies in Europe
CSMS- Combined symptom-medication score
DCP- Digital care pathway
DG- Directorate General
EAACI- European Academy of Allergy and Clinical Immunology
e-DASTHMA- Electronic daily control-medication score in asthma
EtD- Evidence-to-decision
EU- European Union
GARD- Global Alliance against chronic Respiratory Diseases
GRADE- Grading of Recommendations, Assessment, Development and Evaluation
HbA1c- Glycated hemoglobin
ICP- Integrated care pathway
LLM- Large language model
MACVIA- Contre les Maladies Chroniques pour un Vieillessement Actif
MASK- Mobile Airways Sentinel network
OECD- Organisation for Economic Co-operation and Development
PICO- Population, Interventions, Comparators, and Outcomes
RCT- Randomized controlled trial
RWE- Real-world evidence
SDM- Shared decision-making
TLR- Toll-like receptors
UCRAID- Ukrainian Citizen and refugee electronic support in Respiratory diseases, Allergy, Immunology and Dermatology
VAS- Visual analog scale
WHO- World Health Organization

(ARIA), a 2-decade journey, (2) Grading of Recommendations, Assessment, Development and Evaluation (GRADE), the evidence-based model of guidelines in airway diseases, (3)

mHealth impact on airway diseases, (4) From guidelines to digital care pathways, (5) Embedding Planetary Health, (6) Novel classification of rhinitis and asthma, (7) Embedding real-life data with population-based studies, (8) The ARIA-EAACI (European Academy of Allergy and Clinical Immunology) strategy for the management of airway diseases using digital biomarkers, (9) Artificial intelligence, (10) The development of digitally enabled, ARIA person-centered care, and (11) The political agenda. The ultimate goal is to propose ARIA 2024 guidelines centered around the patient to make them more applicable and sustainable. © 2024 The Authors. Published by Elsevier Inc. on behalf of the American Academy of Allergy, Asthma & Immunology. This is an open access article under the CC BY license (<http://creativecommons.org/licenses/by/4.0/>). (J Allergy Clin Immunol Pract 2024;12:2648-68)

Key words: ARIA; Artificial intelligence; Asthma; Evidence-based medicine; Person-centered care; Rhinitis; mHealth

Allergic rhinitis (AR), caused by IgE-mediated reactions to inhaled allergens, is one of the most common chronic conditions globally.¹ AR often occurs concomitantly with asthma and conjunctivitis. AR impairs quality of life; affects social life, school, and work; and is associated with substantial economic costs.^{1,2}

The Allergic Rhinitis and its Impact on Asthma (ARIA) initiative classified AR into intermittent or persistent and mild or moderate/severe, as it proposed guidelines for AR and asthma multimorbidity.³ Over the past 20 years, ARIA has evolved, with strong policymaker commitments, from the first multimorbidity guideline in respiratory diseases³ to GRADE (Grading of Recommendations, Assessment, Development and Evaluation)^{4,5} and next-generation guidelines enhancing the use of patient-centered data (person-centered care and real-world data) and chamber studies.⁶

Applying the GRADE methodology to appraise available evidence has considerably improved the understanding of AR treatment and guideline development.^{4,5,7,8} However, there is an increasing use as well as confusion regarding the role of the so-called real-world evidence (RWE) to inform the clinical practice on concerns about the applicability of the results of randomized controlled trials (RCTs) with restricted inclusion criteria.⁹

^{eeee}Department of Cardiovascular and Respiratory Sciences, Università Cattolica del Sacro Cuore, Rome, Italy

^{ffff}Department of Neurological, ENT and Thoracic Sciences, Fondazione Policlinico Universitario A Gemelli—IRCCS, Rome, Italy

^{gggg}National Heart and Lung Institute (NHLI), Imperial College London, London, United Kingdom

^{hhhh}Allergist, Department of Allergy, La Rochelle, France

ⁱⁱⁱⁱRespiratory Clinic, Department of Internal Medicine, University of Genoa, Genoa, Italy

^{jjjj}Respiratory & Allergy Clinic, IRCCS Ospedale Policlinico San Martino, Genoa, Italy

^{kkkk}Centro Medico Docente La Trinidad, Allergology & Immunology Department, Caracas, Venezuela

^{llll}Department of Pediatrics, Medical School, Federal University of Minas Gerais, Belo Horizonte, Brazil

^{mmmm}NOVA Medical School/Comprehensive Health Research Centre (CHRC), Lisbon, Portugal

ⁿⁿⁿⁿServiço de Imunoalergologia, Centro Hospitalar Universitário de Lisboa Central/ ULS São José, Lisbon, Portugal

^{oooo}Division of Allergy/Immunology, University of South Florida, Tampa, Fla

^{pppp}Department of Respiratory Medicine, Hospital Universitari Dexeus, Barcelona, Spain

^{qqqq}The Group of Rhinitis, Rhinosinusitis and Nasal Polyps, Area of Asthma, SEPAR, Barcelona, Spain

^{rrrr}SOS Allergology and Clinical Immunology, USL Toscana Centro, Prato, Italy

^{ssss}Dr Agostinho Neto University hospital, Epidemiology Department, Praia, Cabo Verde

^{tttt}Immunology, Cabo Verde University, Faculty of Medicine, Praia, Cabo Verde

^{uuuu}Division of Allergy and Clinical Immunology, Seoul National University Bundang Hospital, Seoul National University, College of Medicine, Seoul, Republic of Korea

^{vvvv}Department of Pneumology, Coimbra University Hospital, Faculty of Medicine, Coimbra, Portugal

^{wwww}Coimbra Institute for Clinical and Biomedical Research, CIBB, Coimbra, Portugal

^{xxxx}Faculty of Public Health, Sofia Medical University, Sofia, Bulgaria

^{yyyy}Casa di Cura Villa Montallegro, Allergology Department, Genova, Italy

^{zzzz}Latvian Association of Allergists, University Children Hospital, Riga, Latvia

^{aaaa}Life and Health Sciences Research Institute (ICVS), School of Medicine, University of Minho, Braga, Portugal

Integrated Care Pathways for Airway Diseases (AIRWAYS-ICPs)¹⁰ launched a collaboration to develop multisectoral integrated care pathways (ICPs) for chronic respiratory diseases with a strategic relevance to the European Union Health Strategy and the Digital Single Market. Initiated in 2013 under the frame of the European Innovation Partnership on

Active and Healthy Ageing (Directorate General [DG] Santé and DG Connect),^{10,11} it was a GARD (Global Alliance against chronic Respiratory Diseases, World Health Organization [WHO]) research demonstration project.^{12,13} MASK (Mobile Airways Sentinel network) was developed as the information technology solution¹⁴ to deploy AIRWAYS-ICPs.¹⁵

bbbbb CINTESIS@RISE, Biochemistry Lab, Faculty of Pharmacy and Competence Center on Active and Healthy Ageing, University of Porto, Porto, Portugal
cccc Woolcock Institute of Medical Research, Quality Use of Respiratory Medicine, Sydney, NSW, Australia
ddddd Peercode BV, Geldermalsen, the Netherlands
eeeee Department of Medical Sciences and Public Health and Unit of Allergy and Clinical Immunology, University Hospital "Duilio Casula", University of Cagliari, Cagliari, Italy
fffff VIM Suresnes, UMR 0892, Pôle des Maladies des Voies Respiratoires, Hôpital Foch, Université Paris-Saclay, Suresnes, France
ggggg Medical Faculty Skopje, University Clinic of Pulmology and Allergy, Skopje, Republic of Macedonia
hhhhh Service de Pneumo-Allergologie, Centre Hospitalo-Universitaire de Béni-Messous, Algiers, Algeria
iiiiii Allergy and Clinical Immunology, National Heart and Lung Institute, Imperial College London, London, United Kingdom
jjjjj St Luke's Medical Center, ENT Department, Quezon City, Philippines
kkkkk Allergy, Bambino Gesù Children's Hospital, Istituto di Ricovero e Cura a Carattere Scientifico (IRCCS), Rome, Italy
lllll Department of Otorhinolaryngology, Amsterdam University Medical Centres, AMC, Amsterdam, the Netherlands
mmmmm CHU de Reims, Service des Maladies Respiratoires et Allergiques, Reims, France
nnnnn Department of Internal Diseases, Allergy and Clinical Immunology, Medical University of Silesia in Katowice, Katowice, Poland
ooooo Clínica Ricardo Palma, Allergy & Immunology Department, Lima, Peru
ppppp Department of Internal Medicine 'A. Murri' and Unit of Geriatric Immunology, University of Bari Medical School, Bari, Italy
qqqqq Center of Allergy and Immunology, David Tvildiani Medical University, Tbilisi, Georgia
rrrrr Department of Immunology and Allergy, Lithuanian University of Health Sciences, Kaunas, Lithuania
sssss Immunology and Allergy Division, Clinical Hospital, University of Chile, Santiago, Chile
ttttt Pediatric Allergy, Immunology and Rheumatology Unit, Children's Hospital, Ain Shams University, Cairo, Egypt
uuuuu Department of Clinical Immunology and Allergy, Oncology Institute of St Elisabeth, Bratislava, Slovakia
vvvvv Department of Otorhinolaryngology, Chiba University, Chiba, Japan
wwwww Department of Internal Medicine and Infectious Diseases, St Joseph University, Hotel Dieu de France Hospital, Beirut, Lebanon
xxxxx Department of Allergy and Clinical Immunology, Kazakhstan Association of Allergy and Clinical Immunology, Kazakh National Medical University, Almaty, Kazakhstan
yyyyy Servicio de Alergia e Inmunología, Clínica Santa Isabel, Buenos Aires, Argentina
zzzzz Research Unit of Clinical Medicine, University of Oulu, Oulu, Finland
aaaaa Department of Pediatrics and Adolescent Medicine, University of Turku, Turku University Hospital, Turku, Finland
bbbbb Department of Pulmonology and Phthysiology, Department of Pediatrics, Jessenius Faculty of Medicine in Martin, Comenius University in Bratislava, Bratislava, Slovakia
cccc Institute of Clinical Immunology and Allergy, Jessenius Faculty of Medicine in Martin, Comenius University, Bratislava, Slovakia
ddddd Institute of Clinical Medicine, Children's Clinic, Tartu University, Tartu, Estonia
eeeee Department of Clinical Immunology, Wrocław Medical University, Wrocław, Poland
fffff ALL-MED Medical Research Institute, Department of Clinical Immunology, Wrocław, Poland
ggggg Poltava State Medical University, Immunology & Allergy Department, Poltava, Ukraine

hhhhh Department of Respiratory Medicine, National Institute of Diseases of the Chest and Hospital, Dhaka, Bangladesh
iiiiii Global NCD Platform, Geneva, Switzerland
jjjjj Makerere University Lung Institute, Pulmonology Medicine Department, Kampala, Uganda
kkkkk Department of Otorhinolaryngology, Head and Neck Surgery, Semmelweis University, Budapest, Hungary
lllll Department of Clinical Science and Education, Södersjukhuset, Karolinska Institutet, Stockholm, Sweden
mmmmm Sach's Children and Youth Hospital, Södersjukhuset, Stockholm, Sweden
nnnnn Department of Pediatric Respiratory Diseases and Allergy, Medical University of Warsaw, Warsaw, Poland
ooooo Division of Internal Medicine, Asthma and Allergy, Barlicki University Hospital, Medical University of Lodz, Lodz, Poland
ppppp Department of Clinical and Laboratory Immunology, Allergy and Medical Genetics, Bogomolets National Medical University, Kyiv, Ukraine
qqqqq Institute of Translational Pharmacology (IFT)-National Research Council (CNR), Palermo, Italy
rrrrr Respiratory Medicine, Trinity College, Dublin, Ireland
sssss Respiratory Physician & Allergist, Professorial Respiratory Centre, Tallaght University Hospital & Peamount Healthcare, Dublin, Ireland
ttttt Croatian Pulmonary Society, Clinical Center for Pulmonary Diseases, Zagreb, Croatia
uuuuu Dankook University Hospital, ENT Department, Cheonan, Republic of Korea
vvvvv Asthma, COPD Outpatient Care Unit, University Medical Center, Hô-Chi-Minh City, Vietnam
wwwww Mongolian National University of Medical Sciences, Health Department, Ulan Bator, Mongolia
xxxxx Department of Pulmonary Medicine, CHU Liège, Liège, Belgium
yyyyy GIGA I3 Research Group, University of Liège, Liège, Belgium
zzzzz Rashid Hospital, DUBAI Health, Dubai, UAE
aaaaa Research Institute of Medical and Health Sciences, University of Sharjah, Sharjah, UAE
bbbbb Allergy Unit "D Kalogeromitros", 2nd Department of Dermatology and Venereology, National & Kapodistrian University of Athens, "Attikon" University Hospital, Athens, Greece
cccc Clinic for Pulmonary Diseases, Clinical Center of Serbia, Faculty of Medicine, University of Belgrade, Serbian Association for Asthma and COPD, Belgrade, Serbia
ddddd National Center for Research in Chronic Respiratory Diseases, Collaborating with WHO—EMRO Tishreen University School of Medicine, Latakia, Syria
eeeee Al-Sham Private University, Pharmacy Department, Damascus, Syria
fffff Department of Allergy and Internal Medicine, Medical University of Białystok, Białystok, Poland
ggggg Department of Medicine, Faculty of Medicine and Surgery, University of Malta, Msida, MSD, Malta
hhhhh EPIUnit-Institute of Public Health, University of Porto, and Laboratory for Integrative and Translational Research in Population Health (ITR), Porto, Portugal
iiiiii Serviço de Imunoalergologia, Centro Hospitalar Universitário São João, Porto, Portugal
jjjjj Basic and Clinical Immunology Unit, Department of Pathology, Faculty of Medicine, University of Porto, Porto, Portugal
kkkkk Fundaler, Buenos Aires, Argentina
lllll Rhinology Unit & Smell Clinic, ENT Department, Hospital Clínic, Barcelona, Spain
mmmmm Clinical & Experimental Respiratory Immunology, FRBC-IDIBAPS, CIBERES, University of Barcelona, Barcelona, Spain
nnnnn Université Paris-Saclay, UVSQ, Univ. Paris-Sud, Villejuif, France
ooooo Inserm, Équipe d'Épidémiologie respiratoire intégrative, CESP, Villejuif, France
ppppp Clinical Immunology and Allergy Unit, Sheffield Teaching Hospitals, Sheffield, United Kingdom
qqqqq University of Liverpool, Liverpool, United Kingdom

Digital care pathways (DCPs) employ digital technologies in ICPs.

The traditional healthcare model is focused on diseases (medicine and natural science) and does not acknowledge patients' resources and abilities to be experts in their own lives based on their lived experiences and expectations.^{16,17} Improving healthcare safety, quality, and coordination, as well as health outcomes including quality of life, is an important aim in the care of patients with chronic conditions. Person-centered care

needs to ensure that people's preferences, needs, and values guide clinical decisions. It provides care that is respectful of and responsive to patients and ensures that they are empowered and involved in decision making.¹⁸ Nine themes have been identified in person-centered care: (1) empathy, (2) respect, (3) engagement, (4) relationship, (5) communication, (6) shared decision-making (SDM), (7) holistic focus, (8) individualized focus, and (9) coordinated care.¹⁹ Digital tools are promoters for person-centered care practices in chronic care²⁰ but, alone, cannot

Department of Otolaryngology-Head & Neck Surgery, St Luke's Medical Center, Quezon City, Philippines
Division of Surgery, Philippine Children's Medical Center, Quezon City, Philippines
Center of Allergy, Immunology and Respiratory Diseases, Santa Fe, Argentina
Hungarian Allergy Association, Allergy & Immunology Department, Budapest, Hungary
Department of Allergy, Medical University of Gdańsk, Gdańsk, Poland
Faculty of Medicine, Eduardo Mondlane University, Maputo, Mozambique
ENT Department, University Hospital of Kinshasa, Kinshasa, Democratic Republic of the Congo
Allergy, Asthma and Clinical Immunology, Alfred Health and Department of Immunology, Central Clinical School, Monash University, Melbourne, VIC, Australia
Department of Infection and Immunity, Luxembourg Institute of Health, Esch-sur-Alzette, Luxembourg
Department of Dermatology and Allergy Center, Odense Research Center for Anaphylaxis (ORCA), Odense University Hospital, Odense, Denmark
Japan Antituberculosis Association (JATA), Fukujuji Hospital, Tokyo, Japan
Chiba Rosai Hospital, ENT Department, Chiba, Japan
Chiba University Hospital, Department of Otolaryngology, Head and Neck Surgery, Chiba, Japan
Department of Otolaryngology, Nippon Medical School, Tokyo, Japan
Department of Respiratory Medicine, JSS Medical College, JSSAHER Mysuru, Karnataka, India
Department of Biochemistry and Molecular Biology, School of Chemistry, Complutense University of Madrid, Madrid, Spain
Institute of Pathophysiology and Allergy Research, Medical University Vienna, Vienna, Austria
Interuniversity Messerli Research Institute of the University of Veterinary Medicine and Medical University Vienna, Vienna, Austria
Department of Immunology and Allergy, Faculty of Medicine in Pilsen, Charles University, Prague, Czech Republic
Department of Allergy, University of Helsinki and Hospital for Skin and Allergic Diseases, Helsinki, Finland
Department of Allergy and Clinical Immunology, Ajou University School of Medicine, Suwon, Republic of Korea
Allergy and Respiratory Diseases, IRCCS Policlinico San Martino, University of Genoa, Genoa, Italy
Division of Allergy and Clinical Immunology, Department of Medicine, "Santa Maria della Speranza" Hospital, Battipaglia, Salerno, Italy
Agency of Health ASL, Division of Allergy and Clinical Immunology, Department of Medicine, Salerno, Italy
Postgraduate Programme in Allergy and Clinical Immunology, University of Naples Federico II, Naples, Italy
Department of Pediatrics, Nippon Medical School, Tokyo, Japan
Medical School, University of Cyprus, Nicosia, Cyprus
Prima Nova, Healthcare Institution, Zagreb, Croatia
Medical Faculty, University JJ Strossmayer of Osijek, Osijek, Croatia
Clinic of Occupational Diseases, University Hospital Sveti Ivan Rilski, Sofia, Bulgaria
IRCCS Humanitas Research Center, Personalized Medicine Asthma & Allergy, Rozzano, Milan, Italy
Department of Pulmonology and Allergy, Klaipeda National Hospital, Klaipeda, Lithuania
Vilnius University Medical Faculty, Department of Political Behaviour and Institutions, Vilnius, Lithuania
Division of Adult and Pediatric Allergy and Immunology, University of the Philippines - Philippines General Hospital, Manila, Philippines

Department of Allergy, Clinics Hospital, National University, San Lorenzo, Paraguay
Faculty of Medicine, University of Southampton, Southampton, United Kingdom
The David Hide Asthma and Allergy Centre, St Mary's Hospital, Isle of Wight, United Kingdom
NIHR Southampton Biomedical Research Centre, University Hospital Southampton NHS Foundation Trust, Southampton, United Kingdom
Pneumologie, AP-HP Centre Université de Paris Cité, Hôpital Cochin, Paris, France
UMR 1016, Institut Cochin, Paris, France
Inserm, Epidemiology Department, CESP, Villejuif, France
Pediatric Allergy and Clinical Immunology, Hospital Espanol de Mexico, Mexico City, Mexico
Faculty of Medicine & Surgery, University of Santo Tomas, Manila, Philippines
Division of Allergy, Asthma and Clinical Immunology, Emek Medical Center, Afula, Israel
Rappaport Faculty of Medicine, Technion-Israel Institute of Technology, Haifa, Israel
Department of Otolaryngology-Head and Neck Surgery, Eye and Ear University Hospital, Beirut, Lebanon
Department of Otorhinolaryngology-Head and Neck Surgery, Dar Al Shifa Hospital, Salmiya, Kuwait
ARIA, Asthma, Rhinitis, Immunology & Allergy Department, Athens, Greece
Allergy Service, Fundacion Jimenez Diaz, Autonoma University of Madrid, CIBERES-ISCIII, Madrid, Spain
Asthma Reference Center—School of Medicine of Santa Casa de Misericórdia of Vitória, Espírito Santo, Brazil
SMAIC Société Marocaine d'Allergologie et Immunologie Clinique, Rabat, Morocco
PROMISE Department, University of Palermo, Palermo, Italy
Asthma-Allergy Unit, University Hospital of Verona, Verona, Italy
Clínica Sisul, Allergy & Asthma, Medical Director, CLINICA SISUL, FACAAL, SPAAL, Asuncion, Paraguay
Division of Allergy, Clinical Immunology and Rheumatology, Department of Pediatrics, Federal University of São Paulo, São Paulo, Brazil
Division of Respiratory Medicine, Department of Pediatrics, Hospital Nacional de Niños, Universidad de Costa Rica, San Jose, Costa Rica
Department of Respiratory Medicine and Tuberculosis, University Hospital, Brno, Czech Republic
Head of Laboratory of Atmospheric processes and Aerobiology, University of Latvia, Riga, Latvia
Department of Respiratory Medicine, Copenhagen University Hospital-Hvidovre, Copenhagen, Denmark
Institute of Clinical Medicine, University of Copenhagen, Copenhagen, Denmark
Department of Pediatrics, Victor R Potenciano Medical Center, Mandaluyong, Philippines
Department of Otolaryngology, Faculty of Medicine Siriraj Hospital, Mahidol University, Bangkok, Thailand
Imunolergologia, Centro Hospitalar Universitário de Coimbra, Faculty of Medicine, University of Coimbra, Coimbra, Portugal
Clinic of Allergy and Immunology, University Clinical Center of Serbia, Belgrade, Serbia
Faculty of Medicine, University of Belgrade, Belgrade, Serbia
International Primary Care Respiratory Group IPCRG, Aberdeen, United Kingdom

achieve the ideals of person-centered care.²¹ Politicians and policymakers have an increased interest in adopting and implementing person-centered care.^{22,23}

This global paper (Table E1, available in this article's Online Repository at www.jaci-inpractice.org) reviews the current knowledge to develop (1) DCPs for rhinitis and asthma

multimorbidity and (2) digitally enabled, person-centered care²⁴ using the GRADE approach to integrate both RCTs and RWE. The ultimate goal is to develop guidelines centered around the patient to make them more applicable.

Some diseases have not been considered in this paper (Table I).

Health Planning Unit, Department of Social Medicine, Faculty of Medicine, University of Crete, Iraklion, Greece

Federal University of Pampa, Department of Medicine, Uruguiana, Brazil
Pediatrics, Allergy & Immunology, Latin American Society of Allergy, Asthma & immunology (SLAAi), Montevideo, Uruguay

Department of Lung Diseases and Clinical Immunology, University of Turku, Turku, Finland

Research on Healthcare Performance (RESHAPE), INSERM U1290, Pneumologie et Médecine Respiratoire, Groupement Hospitalier Nord, Hôpital de la Croix Rousse, Lyon, France

FiLHA, Finnish Lung Health Association, Helsinki, Finland

Department of Clinical Medicine, Pulmonary Diseases and Clinical Allergology, University of Turku, Turku, Finland

Division of Allergy and Immunology, Department of Pediatrics, Siriraj Hospital, Mahidol University Faculty of Medicine, Bangkok, Thailand

Pulmonary Environmental Epidemiology Unit, CNR Institute of Clinical Physiology, Pisa, Italy

Nova Southeastern University, College of Allopathic Medicine, Fort Lauderdale, Fla

Department of Otolaryngology, Yong Loo Lin School of Medicine, National University of Singapore, Singapore, Republic of Singapore

Department of Medicine, Clinical Immunology and Allergy, McMaster University, Hamilton, ON, Canada

Department of Paediatrics, Prince of Wales Hospital, Hong Kong, SAR, China

Division of Allergy and Immunology, Department of Dermatology, Allergy and Venerology, Charité Universitätsmedizin Berlin, Berlin, Germany

The Allergy and Asthma Institute, Allergy & Asthma Department, Islamabad, Pakistan

Clemenceau Medical Center DHCC, Lebanese-American University, Dubai, UAE

University Clinic of Respiratory and Allergic Diseases, Pulmonary & Allergy Department, Golnik, Slovenia

Faculty of Medicine, University of Ljubljana, Ljubljana, Slovenia

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ARIA: A 2-DECADE JOURNEY

ARIA was initiated during a WHO workshop in 1999.³ It has evolved in 6 phases.

Phase 1 (1999-2009)

- Development and update of an evidence-based document³⁷ to provide a guide for the diagnosis and management of AR and asthma multimorbidity by physicians^{3,38} and pharmacists.³⁹ A specific focus was placed on developing countries.

personal fees from Laboratorios Casasco Argentina, outside the submitted work. J. Sastre reports grants and personal fees from Sanofi, personal fees from GSK, personal fees from Novartis, personal fees from AstraZeneca, personal fees from Mundipharma, and personal fees from Faes Farma, outside the submitted work. S. Quirce reports personal fees and nonfinancial support from GSK, personal fees and nonfinancial support from AstraZeneca, personal fees and nonfinancial support from Sanofi, personal fees and nonfinancial support from Novartis, personal fees and nonfinancial support from Mundipharma, personal fees and nonfinancial support from Allergy Therapeutics, and personal fees and nonfinancial support from Chiesi, outside the submitted work. S. Toppila-Salmi reports personal fees from AstraZeneca, personal fees from ALK-Abelló, personal fees from GSK, personal fees from Sanofi, personal fees from Clario, personal fees from Orion Pharma, grants from GSK, and grants from Sanofi, outside the submitted work. O. Palomares reports fees for Lectures/Advisory Boards from AstraZeneca, Pfizer, GSK, Immunotek SL, Novartis, Sanofi-Genzyme, and Regeneron. He has received research grants from MINECO, MICINN, CAM, Immunotek SL, Novartis, and AstraZeneca. M. Ollert reports personal fees from Hycor Diagnostics, personal fees from Allergy Therapeutics/Bencard, outside the submitted work; and is the scientific co-founder of Tolerogencis SARL, Luxembourg. V. Kvedariene reports nonfinancial support from Norameda, nonfinancial support from Berlin Chemie Menarini, and nonfinancial support from Dimuna, outside the submitted work. A. Todo-Bom reports personal fees from GSK (GlaxoSmithKline), personal fees from AstraZeneca, nonfinancial support from Leti, nonfinancial support from Abbvie, and personal fees from Viatriis, outside the submitted work. M. Kupczyk reports personal fees from Adamed, personal fees from AstraZeneca, personal fees from Berlin Chemie Menarini, personal fees from Chiesi, personal fees from Celon Pharma, personal fees from GSK, personal fees from Novartis, personal fees from Sanofi Aventis, personal fees from Abbvie, personal fees from Aurovitas, personal fees from LEK-AM, personal fees from Teva, personal fees from Zentiva, personal fees from Polpharma, personal fees from HAL-Allergy, personal fees from Emma, and personal fees from Opella Healthcare, outside the submitted work. O. Pfaar reports grants and personal fees from ALK-Abelló, grants and personal fees from Allergopharma, grants and personal fees from Stallergenes Greer, grants and personal fees from HAL Allergy Holding BV/HAL Allergie GmbH, grants from Bencard Allergie GmbH/Allergy Therapeutics, grants from Lofarma, grants and personal fees from ASIT Biotech Tools S.A., grants and personal fees from Laboratorios LETI/LETI Pharma, grants and personal fees from GlaxoSmithKline, personal fees from ROXALL Medizin, personal fees from Novartis, grants and personal fees from Sanofi-Aventis and Sanofi-Genzyme, personal fees from Med Update Europe GmbH, personal fees from streamedup! GmbH, grants from Pohl-Boskamp, grants from Immunotek S.L., personal fees from John Wiley and Sons, AS, personal fees from Paul-Martini-Stiftung (PMS), personal fees from Regeneron Pharmaceuticals Inc., personal fees from RG Aerztefortbildung, personal fees from Institut für Disease Management, personal fees from Springer GmbH, grants and personal fees from AstraZeneca, personal fees from IQVIA Commercial, personal fees from Ingress Health, personal fees from Wort&Bild Verlag, personal fees from Verlag ME, personal fees from Procter&Gamble, personal fees from ALTAMIRA, personal fees from Meinhardt Congress GmbH, personal fees from Deutsche Forschungsgemeinschaft, personal fees from Thieme, grants from Deutsche AllergieLiga e.V., personal fees from AeDA, personal fees from Alfred-Krupp Krankenhaus, personal fees from Red Maple Trials Inc., personal fees from Königlich Dänisches Generalkonsulat, personal fees from Medizinische Hochschule Hannover, personal fees from ECM Expro&Conference Management, personal fees from Technical University Dresden, personal fees from Lilly, personal fees from Paul Ehrlich Institut, personal fees from Japanese Society of Allergy, personal fees from Forum für Medizinische Fortbildung, from Dustri-Verlag, outside the submitted work. He is the member of EAACI Excom; the member of the external board of directors of DGAKI; coordinator, main author, or

- Dissemination and implementation: ARIA has been translated into over 50 languages, and disseminated and implemented in over 80 countries.⁴⁰
- Update using the same evidence-based system.^{37,38}

Phase 2 (2010-2016)

- ARIA was revised using the GRADE approach for assessing the strength of evidence underpinning recommendations. ARIA was one of the first guidelines to use GRADE evidence-

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to-decision (EtD) frameworks.^{4,5} An update was published in 2017.⁵

- Deployment to policymakers.⁴⁰

Phase 3 (2016-2018)

- An algorithm (MACVIA: Contre les Maladies Chroniques pour un Vieillissement Actif) was devised⁴¹ and digitalized⁴² to step up or step down AR treatment based on control (Figures 1 and 2).⁴¹ Algorithms require testing with RWE that includes RCTs and observational research with person-centered care.⁴³⁻⁴⁵ A consensus refined the algorithm.⁴¹
- Implementation of mHealth tools for individualized and predictive medicine to develop ICPs for the management of AR and asthma by a multidisciplinary team centered on the needs of patients (MASK).⁴⁶⁻⁴⁸
- Initiation of MASK-air.⁴⁶

Phase 4 (2018-2019)

- Digital transformation of health and care.²⁴
- Change management to improve population health and provide well-being for rhinitis and asthma sufferers across the life cycle, irrespective of their gender, age, or socioeconomic status and with the overarching aim to reduce health and social inequities.⁴⁹
- Development of MASK-air, the ARIA app.

Phase 5 (2019-2021)

- Next-generation guidelines for the AR pharmacologic treatment were developed using existing GRADE-based guidelines for the disease, RWE provided by mobile technology, and additive studies (allergen chamber studies) to refine the MACVIA algorithm.⁶
- MASK-air, a Good Practice of DG Health and Food Safety: Digitally enabled, patient-centered care.^{24,50}
- Value-added medicine for the repurposing of AR medications.⁵¹

- High-level meeting (Finnish Presidency of the European Union [EU]) on Planetary Health.⁵²

Phase 6 (2022-)

- Participation in initiatives on Planetary Health with climate change (EU Horizon Europe grant CATALYSE [Climate Action To Advance HealthY Societies in Europe] 2022-7).⁵³
- Digitally enabled, person-centered care including next-generation care pathways embedding RWE based on data with the GRADE evaluation of interventions.

ARIA has participated in several EU or WHO projects and grants, and European Academy of Allergy and Clinical Immunology (EAACI) Task Forces (Table E1, available in this article's Online Repository at www.jaci-inpractice.org).

GRADE, THE EVIDENCE-BASED MODEL OF GUIDELINES IN AIRWAY DISEASES

Strengths

To evaluate the confidence in the evidence underlying estimates of effects of interventions and to develop recommendations in guidelines, the GRADE^{4,5} methodology explicitly considers all types of study designs from RCTs to case reports, although guideline developers often restrict guidelines to RCTs.⁵⁴⁻⁵⁶ For the formulation of recommendations on interventions, GRADE considers not only their benefits and harms, but also—and among others—patients' values and preferences, costs and cost-effectiveness, acceptability, and feasibility. The EtD of GRADE allows the evidence to be considered on all of these criteria, based on which recommendations are formulated.

The ARIA revision 2016,⁵ the US Practice Parameters 2017,⁷ and 3 questions of the US Practice Parameters 2020⁸ used GRADE as their methodological approach. Interestingly, the same questions were considered, and the results of these guidelines supported the MACVIA algorithm.⁴¹ The ARIA revision

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TABLE I. Excluded diseases

Although nonallergic rhinitis (NAR) is very common and may be associated with allergic rhinitis, it cannot be considered in Allergic Rhinitis and its Impact on Asthma (ARIA) because (1) there are many distinct NAR diseases²⁵ and many phenotypes that may overlap and are still poorly defined.²⁶ (2) Although in clinical practice, questionnaires²⁷ and treatments are proposed,²⁸ many medications have been tested in randomized controlled trials (RCTs), and most were not effective, possibly because the NAR phenotypes were not characterized.²⁹ Moreover, many trials have not been published because of lack of efficacy. In published trials, the effect of treatment is often insignificant,³⁰ low, incomplete, or found only in some types of NAR.^{31–34} Observational studies cannot be used in guideline development if NAR phenotypes are not considered. Well-conducted RCTs in different phenotypes of NAR are required to further advance our understanding of the effectiveness of treatments in NAR. Based on these limitations, a meta-analysis would be difficult to interpret and recommendations for NAR cannot be developed using the EtD (evidence-to-decision) Grading of Recommendations, Assessment, Development and Evaluation (GRADE) method used in this document. **Local allergy** is a well-characterized phenotype,³⁵ and it was identified as a research question by the ARIA group. However, it was not prioritized because there is apparently no pharmacologic RCT in this IgE-mediated phenotype.³⁶ Large-scale observational studies are also lacking.

was used as the case scenario on the review published on “How to interpret guidelines.”⁵⁷

In cluster-randomized trials, guideline-driven treatment was reported to be more effective than free treatment choice.^{58,59} Moreover, guidelines (in AR or asthma) have led to a better understanding of the treatment of the disease and have had an important teaching role that has led to a change in the management.⁴⁹ Evidence from direct patient data, however, suggests that guidelines are not sufficiently followed, possibly because they would need to be closer to patients’ concerns.

Combining information from RCTs with real-world data studies

The applicability of the results of RCTs is restricted because of some serious issues (Table II).⁹

There is an increasing trend to use person-centered care to inform the clinical practice, especially as RCTs are often limited to the generalizability and applicability of results.⁶¹ The trade-off that is made is one between risk of bias, primarily selection and confounding bias, and applicability. Ideally, both types of evidence are merged in a way to reduce bias and increase applicability.⁹

pure subpopulations of white blood cells in immune system disorders pending. Y. Okamoto reports personal fees from Torii Pharmaceutical Co., Ltd, personal fees from Tanabe-Mitsubishi Pharmaceutical Co., Ltd, personal fees from Kirin Holdings Co., Ltd, personal fees from Novartis Co., Ltd, personal fees from Allergologisk Laboratorium København, personal fees from Shionogi Co., Ltd, personal fees from Stallergenes-Greer, and personal fees from Daiichi-Sankyo, outside the submitted work. B. Cvetkovski reports personal fees from GSK Pty Ltd, personal fees from Viatris, and personal fees from Sanofi, outside the submitted work. F. M. Tan reports personal fees from Intermed, Pediatrca, A. Menarini, Nestle, and Cathay Drug: honoraria for lectures/presentations and/or module development; and is the board member (Secretary) of the Philippine Society of Allergy, Asthma and Immunology. The rest of the authors declare that they have no relevant conflicts of interest.

THE mHealth IMPACT ON AIRWAY DISEASES

mHealth in allergic diseases and asthma

To select apps for rhinitis, a new approach to market research was based on the automatic screening of the Apple App Store and Google Play Store using JavaScript.⁶² Three apps were available internationally and were being used in 2021 (Vienna Pollen,⁶³ AllergyMonitor,^{64,65} and MASK-air⁶⁶).

MASK-air was developed to implement AIRWAYS-ICPs. It is an app centered around the patient⁴⁸ and is operational in 27 countries and 19 languages (Table E1). Approximately 35,000 users with AR and/or asthma have been registered. MASK-air has been classified as a Medical Device Regulation Class IIa. It is a good practice of DG Santé on digitally enabled, patient-centered care.²⁴ It is also a best practice of OECD (Organisation for Economic Co-operation and Development). MASK-air data have enabled large observational person-centered care studies, novel phenotype discovery, and characterization,⁶⁷ as well as novel insights into the management of AR.^{68–70} MASK-air has also allowed for the development and validation of the ARIA-EAACI combined symptom-medication score for allergic diseases (CSMS) and a daily electronic asthma symptom-medication score (e-DASTHMA).⁷¹

Messages from MASK-air in rhinitis pertinent to guideline development

Several digital studies in up to 35,000 users (39,000 weeks with 6 or 7 days of reporting and over 5000 months with over 26 days of reporting) in 27 countries enabled an assessment of AR treatments.⁷² Their results yield important observations that should be considered in the management of AR (Table III).

The current medications for AR are centered around continuous long-term treatment, and medication registration is based on RCTs carried out for a minimum of 14 days with adherence $\geq 70\%$. Similar to the Global Initiative of Asthma in asthma, a novel approach to treating AR involves suggesting an as-needed treatment regimen based on the presence and severity of symptoms, as opposed to the traditional continuous treatment approach.⁵¹

Digital health in shared decision-making

There is a complete disconnection between the physician’s prescription and the patient’s behavior for the treatment of pollen-induced AR.⁶⁸ The vast majority of allergists prescribe medications for the entire season, recommending the patient to use them regularly, even on days with few symptoms. Some allergists prescribe a preseason treatment without clear evidence of efficacy.⁷⁹ On the other hand, the vast majority of patients use

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Corresponding author: Jean Bousquet, MD, Institute of Allergology, Charité—Universitätsmedizin Berlin, Campus Benjamin Franklin, Hindenburgdamm 30, Haus II, Berlin 12203, Germany. E-mail: jean.bousquet@orange.fr.

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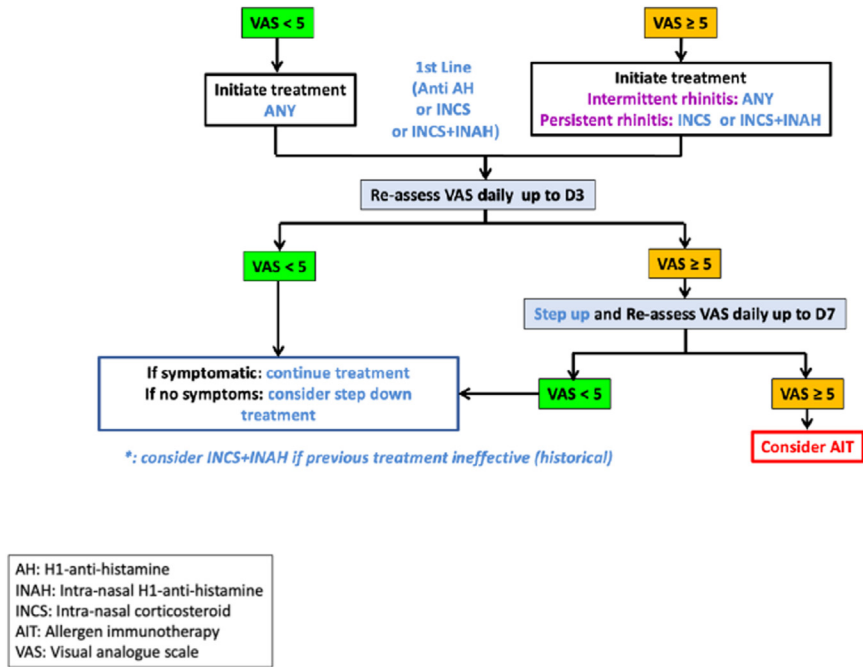


FIGURE 1. Management algorithm of untreated symptomatic patients using control (VAS). Reprinted with permission from Bousquet et al.⁴¹

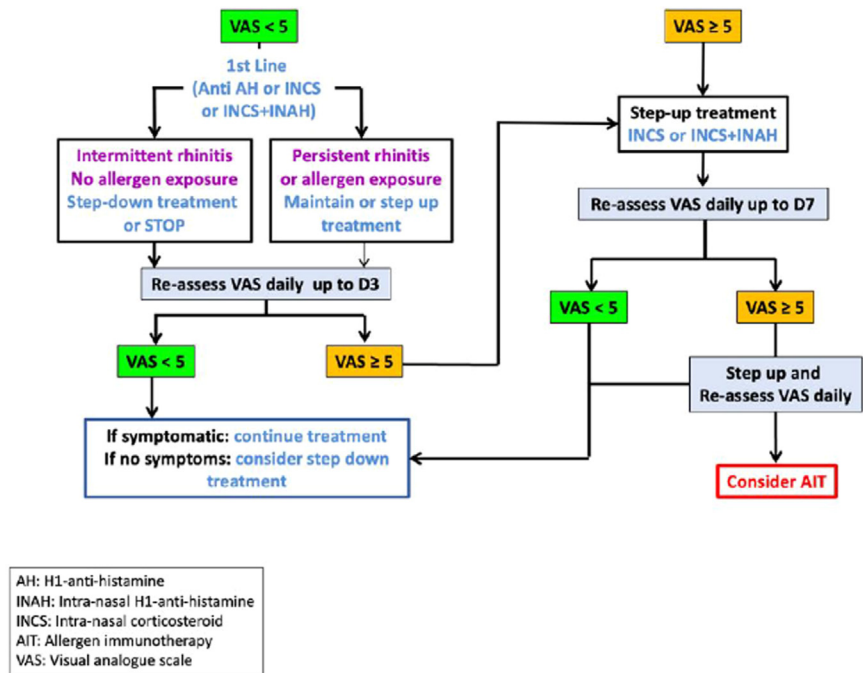


FIGURE 2. Management algorithm of treated symptomatic patients using control (VAS). Reprinted with permission from Bousquet et al.⁴¹

their medications on demand, when their AR is not well controlled. They do not follow the guidelines.^{47,48} When physicians are patients themselves, they behave like patients when they treat their own AR and do not follow the prescriptions they would usually recommend to their patients.⁸⁰ Health literacy is an important component of adherence to medications,^{81,82} but, given the behavior of allergists as patients,

it appears that other factors are also important. Human behavior appears to be a major driver of adherence. The shift from a paternalistic model of health care to a doctor-patient relationship (in which the doctor and patient make shared decisions [SDM]) requires an actively involved patient who takes responsibilities.^{83,84} Rather than being passive, mHealth solutions provide the opportunity for the patient to be

TABLE II. Some weaknesses of randomized controlled trials (RCTs) of rhinitis interventions

	RCT	Putative problems
Severity/control	The worst controlled patients	The recommendations may not apply to patients with mild or partly controlled symptoms that represent the largest population of patients
Patient selection criteria	Asthma usually excluded	• Treatment differences exist in patients with rhinitis alone or rhinitis and asthma • However, patients with uncontrolled disease may respond equally • Do not consider the patient's experience with previous treatments
	Other exclusion criteria	Less than 10% of patients seen in primary care can be enrolled in RCTs ⁶⁰
Adherence to treatment	In most studies $\geq 70\%$	Only a small subset of patients with rhinitis is adherent $\geq 70\%$

TABLE III. Patient-centered lessons in rhinitis provided by MASK-air data

• Patients are poorly adherent to treatment.^{73,74} • Most patients with allergic rhinitis use on-demand treatment when they are suboptimally controlled. This is suggested by the fact that days on which patients do not take medications are usually well controlled.^{68,72,73,75,76} Switching of treatment is common.^{68,75} • The vast majority of patients do not follow the prescriptions of their physicians, who, often, do not follow guidelines.⁷²⁻⁷⁴ Medication use peaked during the pollen season in all of the investigated European countries,⁷⁶ whereas cultural behaviors, assessed using Google Trends,⁷⁷ differed. Oral antihistamines (OAH) were the most common medications reported in monotherapy and combined medications (comedication). This is against guideline recommendations and does not accord with the dispensing of medications (over-the-counter and prescribed) in the pharmacy.⁷⁷ • On most of the days with patients reporting worse control, an increased number of medications are used.^{68-70,72,73} This accords with the concept of Severe Chronic Upper Airway Disease.⁷⁸ • Days with OAH monotherapy are associated with a poorer level of control than days with intranasal corticosteroid (INCS)-containing medications. Days with INCS are associated with a poorer control than those with azelastine-fluticasone (MPAzeFlu).^{72,73} Days with comedication use are associated with a poorer level of control than those reporting monotherapy.^{68-70,72,73}

an active participant in his or her health.^{85,86} Informed self-management is a crucial aspect of patient care in AR, but, as evidenced by MASK-air, most patients do not adhere to the recommended treatment regimens.⁶⁸

USE OF ARTIFICIAL INTELLIGENCE IN GUIDELINE DEVELOPMENT

In the future, artificial intelligence (AI) is most likely to have an important impact on guideline development. Currently, it is already adept at expediting the evidence synthesis and translating content between languages, but also generating plain language summaries. If used appropriately (eg, using Retrieval Augmented Generation), it can help patients navigate contents of the guidelines.

ARIA 2024 will use ChatGPT in two different ways for question generation: (i) we will prompt ChatGPT to either assume the role of a patient or of a healthcare provider and provide relevant guideline questions in the

PICO format; (ii) we will retrieve popular queries on allergic rhinitis using Google Trends and use ChatGPT to classify these queries into those conveying potentially relevant questions versus those not conveying questions (queries identified as potentially conveying relevant questions will then be manually transformed into guideline questions in the PICO format).

In the future, AI-based methods may be used to support the analysis of real-world data (including direct patient data from MASK-air), allowing us to obtain findings that may support the development of guideline recommendations.

PATIENT VALUES AND PREFERENCES

Healthcare interventions typically result in benefits and harms. Patients' values and preferences concern the relative importance patients place on specific benefits and harms. Taking them into account is therefore essential for patient-centered guidelines. For example, in AR, antihistamines can lead to reduced allergy symptoms (benefits) and to an increased risk of side effects (risks). In order to formulate recommendations on antihistamines, we would need to consider the importance that patients attribute to the possibility of having their nasal symptoms improved over the risk of having mild side effects. Values and preferences can be quantitatively measured using different approaches, the most common of which involves utilities. Given that values and preferences are one of the criteria of the GRADE EtD framework, they will be considered in the context of the ARIA guidelines. In fact, a systematic review of patients' values and preferences for health states in AR has been conducted.⁸⁷ The results of this systematic review will be considered when judging the balance of effects when comparing different interventions. For example, in the comparison between 2 equally safe treatments, we will say that a treatment that results in a greater improvement in nasal symptoms may be favored over another that results in a greater improvement in ocular symptoms (as patients tend more often to rate nasal symptoms as more important than ocular symptoms).

FROM GUIDELINES TO DCPs

ICPs are structured multidisciplinary care plans detailing the key steps of patient care.⁸⁸ They promote the translation of guideline recommendations into local protocols and their application to the clinical practice. They may be of particular interest in patients with multimorbidities because guidelines often fail to adequately address their specific needs and

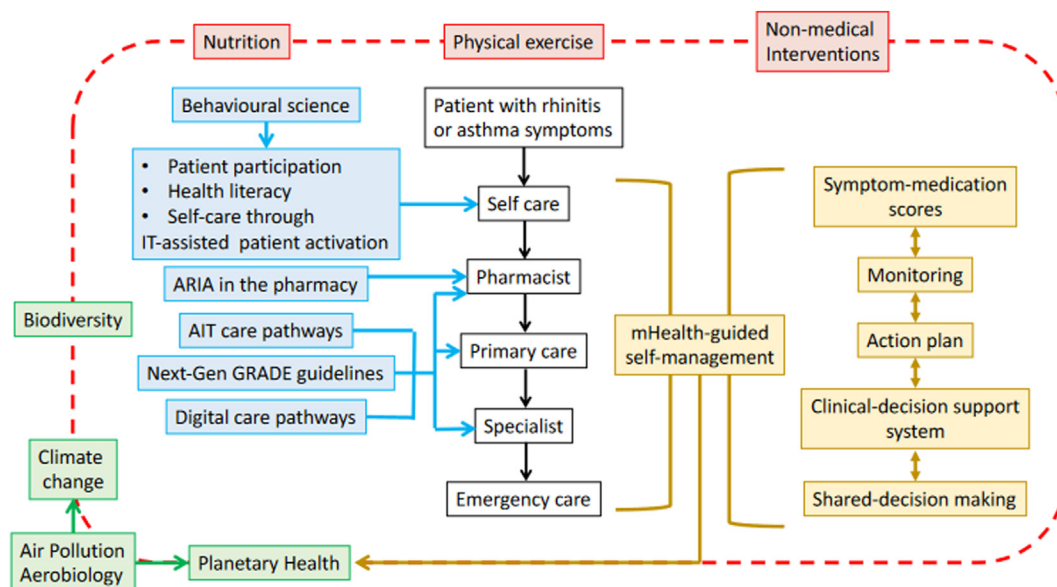


FIGURE 3. Digital care pathways in rhinitis and asthma and the evidence ecosystem. *AIT*, Allergen immunotherapy; *ARIA*, Allergic Rhinitis and its Impact on Asthma; *GRADE*: Grading of Recommendations, Assessment, Development and Evaluation. Reprinted with permission from Bousquet et al.¹

concerns.^{89,90} ICPs should be carried out by a multidisciplinary team including physicians, pharmacists,^{91,92} and allied healthcare professionals.⁹³ ICPs should integrate recommendations from clinical practice guidelines, but they usually (1) enhance recommendations by combining interventions, integrating quality assurance, and (2) describe care coordination. Self-care and SDM are at the forefront of ICPs with the aim of empowering patients and their (professional/lay) caregivers.

DCPs should incorporate all the steps of disease management in a multisectoral ICP using digital technologies. In ARIA 2019-2022, several consensus documents were produced for ICPs:^{15,94,95} ARIA in the pharmacy,⁹⁶⁻⁹⁸ allergen immunotherapy (AIT),^{99,100} and next-generation guidelines.⁶ However, DCPs need to embed environmental triggers^{101,102} and extend their recommendations to nonmedical treatments.¹⁰³ Indoor and outdoor pollution are important to include, but it is not known whether air pollution increases the severity of AR and/or its prevalence.¹⁰⁴⁻¹⁰⁶ Biodiversity, climate change,¹⁰⁷ and Planetary Health should also be considered (Figure 3).^{52,108-110}

EMBEDDING PLANETARY HEALTH AND NATURE DEFICIENCY IN THE ARIA FRAMEWORK

There is an urgent need to safeguard our planet and our health in line with the Declaration of Helsinki.^{109,111} To protect human health in the Anthropocene epoch, human health and the health of the Planet should go together.^{112,113} In AR, as in other chronic diseases, it is important to understand both its close connection to natural systems and how much AR care affects the health of the Planet. *Nature (biodiversity) loss* is the loss or decline of the state of nature.¹¹⁴ A novel concept, *nature deficiency*, refers to nature loss in the human body influencing health. The urban-like environment and lifestyle have weakened the connection of the human body as an ecosystem to wider ecosystems.

ARIA has already been involved in these actions that now need to be deployed to citizens. During the high-level meeting (Finnish Presidency of the EU and DG Research) on Planetary Health,^{109,111} there was a session on MASK-air in the frame of Impact of air POLLution on Asthma and Rhinitis (POLLAR).⁵² MASK-air is one of the partners of a new Horizon Europe grant, CATALYSE (grant agreement number 101057131).⁵³ One of the CATALYSE aims is to develop early warning systems and predictive models to improve the effectiveness of adaptation strategies to climate change, including a specific tool for AR.

The ARIA 2024 guidelines will attempt to embed considerations of Planetary Health into guideline development by including it in the EtD framework for the formulation of recommendations. When producing guideline recommendations, one aspect that will be taken into account will be how the interventions fare in terms of their impact on planetary health. For example, for comparisons between intranasal versus oral treatments, aspects such as the global warming potential and ozone depletion potential of the different packaging types will be assessed. However, while we begin piloting this, the methodological approaches necessary to embed Planetary Health into guidelines are under development.¹¹⁰

NOVEL CLASSIFICATION OF RHINITIS AND ASTHMA

Allergic diseases (asthma, AR, and atopic dermatitis in early life) are associated with allergen-specific IgE and nonallergic mechanisms that may coexist. These diseases tend to cluster, and patients present concomitant or consecutive diseases (multimorbidity). Substantial clinical and immunological differences exist between mono- and polysensitized subjects.^{115,116} The concept of “one-airway-one-disease,” coined over 20 years ago,³ is a simplistic approach of the links between upper- and lower-airway allergic diseases.¹¹⁷ Moreover:

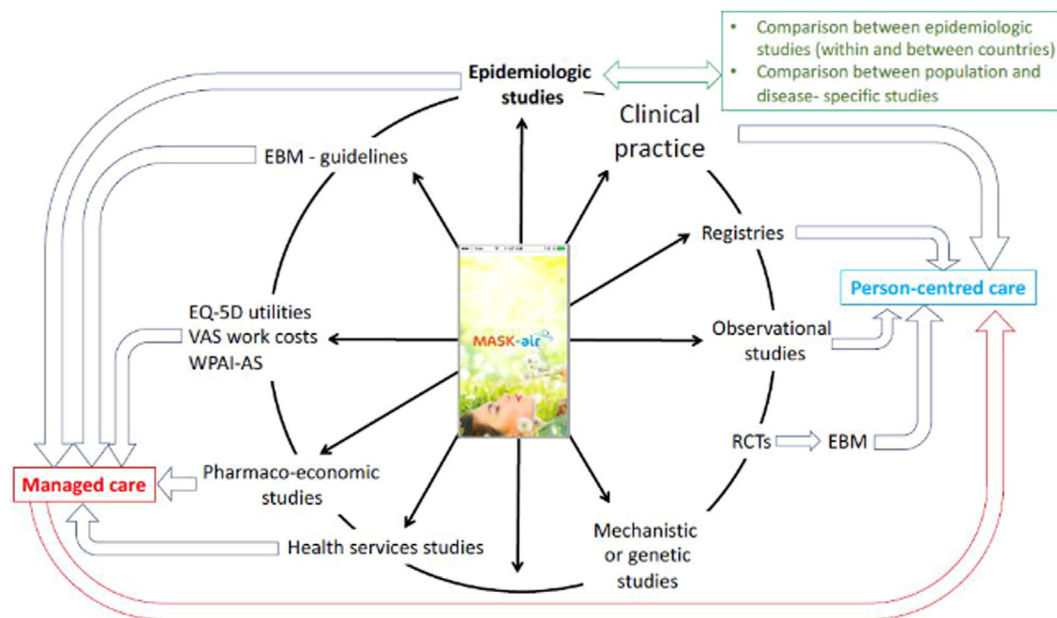


FIGURE 4. Impact of the interaction between MASK-air and population studies. *EBM*, evidence-based medicine; *MASK*, Mobile Airways Sentinel network; *RCT*, randomized controlled trial; *VAS*, visual analog scale; *WPAI-AS*, work productivity and activity impairment allergic specific.

- The clinical observations that led to ARIA clearly indicated that only 30% of patients with rhinitis have asthma, whereas most patients with asthma have rhinitis.^{118,119}
- In birth and children's cohorts, mono- and polysensitization to different allergens represent expressions of distinct diseases.^{120,121} Compared with monosensitization, polysensitization was linked to more robust global IgE response, disease phenotypes (rhinitis alone vs asthma+rhinitis), symptoms, and trajectories. Multimorbidity is partly independent of IgE sensitization, suggesting distinct causal (genomic and mechanistic) pathways.¹²² There is an association between IgE polysensitization and multimorbidity including age of onset, number of allergic multimorbidities (conjunctivitis and atopic dermatitis), severity of disease,¹²³ eosinophil levels, and total IgE levels.
- The MASK-air study showed that there is a multimorbid phenotype (asthma+rhinitis+conjunctivitis) associated with more severe symptoms and a higher impact of symptoms on work productivity compared with the observations with individual diseases.⁶⁷ This phenotype was confirmed using rhinitis⁷⁵ or asthma¹²⁴ to perform cluster analyses.
- These data were confirmed in canonical epidemiologic studies.^{125,126} Rhinitis and rhinoconjunctivitis are separate diseases. The extreme allergy phenotype including asthma+rhinitis+conjunctivitis has been confirmed.¹²⁷⁻¹³³ For all parameters studied, multimorbidity differs from asthma or rhinitis alone. In the French general population epidemiologic study, Constances, participants with asthma+rhinitis had more severe symptoms than those with rhinitis alone as well as an earlier age of onset.¹³³ This suggests that multimorbidity behaves differently than rhinitis alone.
- Genomic findings: Two methods (transcriptomics and RNA sequencing) yielded the same results in 2 different cohorts (Mechanisms of the Development of Allergy and Epigenetic Variation and Childhood Asthma in Puerto Ricans):

multimorbidity was associated with 7 genes of T2 signaling: *IL5* (eosinophils) and *IL33* (polysensitization and eosinophilia).¹³⁴ A total of 27 genes were identified for rhinitis alone and included Toll-like receptors (TLR) and IL-17. These studies suggest that rhinitis alone is a local IL-17-driven disease, whereas T2-associated rhinitis+asthma are systemic IL-33-driven diseases. There are shared epigenetic patterns of allergic multimorbidities, but, in children, these patterns were found only in rhinitis+asthma (and not in asthma alone).

- There are therapeutic differences between patients with rhinitis and patients with rhinitis+asthma. Multimorbid patients more often reported a treatment with intranasal corticosteroids and oral antihistamines,¹³³ which is associated with poor control.⁷⁰ In MASK-air, the comedication pattern was associated with a poorer rhinitis control than in monotherapy.^{68,72} In the CSMS, the distinction between rhinitis and asthma+rhinitis was clear with large effect sizes.

These studies lead to the recognition of 2 distinct diseases: rhinitis alone (local, IL-17, and TLR associated) and rhinitis+asthma (systemic and IL-33 associated) with almost no overlap.¹¹⁷ This new classification needs to be integrated in guideline development, namely by providing—whenever justified—recommendations for patients with rhinitis alone versus rhinitis with asthma.

REAL-LIFE DATA FROM POPULATION-BASED STUDIES

Embedding MASK-air data from general population studies allows the bridging of several fields to assess the relevance of RCTs, observational studies, registries, research in the general population, and others. It appears to be particularly important to compare population and disease-specific epidemiologic studies as an essential step for person-centered care (Figure 4).

TABLE IV. Potential implications of the allergy combined symptom-medication score (CSMS)

1 <i>Clinical practice</i>
• Indication of a treatment in stratified patients
• Follow-up of a treatment and early stopping rule
• Follow-up of a treatment and regular review of efficacy
• Follow-up of the patient when the treatment is stopped
• Reintroduction and follow-up of the treatment in patients who relapsed
2 <i>Randomized controlled trials (RCTs)</i> : mHealth biomarkers are currently exploratory end points but may become primary end points mimicking real life after validation.
3 <i>Observational studies</i> can triangulate RCTs and make a link with the clinical practice.
4 <i>Direct-patient data (real-world data)</i> are the data relating to patient health status and/or the delivery of health care routinely collected from a variety of sources including apps. These data can be obtained by performing large simple trials and pragmatic clinical trials.
5 <i>Epidemiologic studies</i> will use the same approach to better relate RCTs and the clinical practice.
6 <i>Allergen challenge</i> can triangulate RCTs and make a link with the clinical practice.

From Bousquet et al.¹³⁵

**THE ARIA-EAACI STRATEGY FOR THE
MANAGEMENT OF AIRWAY DISEASES USING
DIGITAL BIOMARKERS**

Biomarkers for the diagnosis, treatment, and follow-up of patients with asthma or rhinitis are urgently needed. Although some biologic biomarkers exist in specialist care for asthma (eg, sputum eosinophils or fractional exhaled nitric oxide), they cannot be largely used in primary care. There are no validated biomarkers in rhinitis or AIT that can be used in the clinical practice. The digital transformation of health and health care (including mHealth) places the patient at the center of the health system and is likely to optimize the practice of allergy. ARIA and EAACI developed a task force aimed at proposing digital biomarkers that can be easily used for different purposes in AR and asthma and that form a bridge between the clinical practice, RCTs, and allergen challenges.¹³⁵ Using the MASK-air app as a model, a daily electronic CSMS for allergic diseases¹³⁶ and asthma (e-DASTHMA)⁷¹ was embedded in a strategy similar to the diabetes approach for disease control. The potential implications for the management of allergic respiratory diseases were proposed (Table IV).

In diabetes, 2 types of biomarkers are defined to monitor disease control.^{137,138} The daily control monitoring is assessed using glycemia measurement and longer-term monitoring using glycated hemoglobin (HbA1c) measurement. It is recommended that both tests should be used to optimize diabetes management. By analogy with the diabetes approach, 2 types of patient-centered digital biomarkers are available for rhinitis and asthma:

- Long-term monitoring using control scores (analogous to HbA1c measurement): CARAT (Control of Allergic Rhinitis and Asthma Test)¹³⁹⁻¹⁴¹ is proposed as it combines rhinitis and asthma control. Furthermore, there is a recall period of 4 weeks, whereas many other rhinitis (eg, Allergic Rhinitis Control Test¹⁴² and Rhinitis Control Assessment Test¹⁴³) or asthma (eg, Asthma Control Questionnaire¹⁴⁴) control questionnaires are based on a 1-week recall period. The Asthma

Control Test is based on a 4-week period.¹⁴⁵ These questionnaires, however, do not fully capture the control in patients with fluctuating symptoms (particularly those with severe asthma).

- Daily monitoring of the control (analogous to glycemia measurement): This can be measured using the ARIA-EAACI allergy CSMS¹³⁶ or the e-DASTHMA.⁷¹

**DEVELOPMENT OF DIGITALLY ENABLED, ARIA
PERSON-CENTERED CARE**

The development of guidelines according to the GRADE methodology involves a stepwise approach resulting in the formulation of recommendations for a set of selected questions. For ARIA, we propose the development of guidelines that are (1) digitally enabled, by formally integrating into the guideline development process real-life data obtained from mobile apps such as MASK-air and web searches; (2) person-centered, by taking into account patients' values and preferences when issuing recommendations (as recommended by GRADE); and (3) AI-assisted, by formally integrating large language models (LLMs) into the guideline development process (Figure 5).

**Generation and prioritization of PICO questions
(step 1)**

Question generation. Questions for the ARIA guidelines will follow the Population, Interventions, Comparators, and Outcomes (PICO) framework. In the first phase, questions to be considered by the panel members will be:

- (1) Questions developed for ARIA 2010⁴ and 2016.⁵
- (2) Questions suggested by panel members: Panel members will suggest questions that have not been considered in ARIA 2010 or 2016, which may include some of the questions of the US practice parameters.⁸
- (3) Real-life data-driven questions: Studies based on MASK-air data will be systematically assessed by 2 independent members of the methodology team. In addition, AR-related popular queries will be obtained using Google Trends, as web searches may provide a glimpse into what is of most interest to internet users and, therefore, contribute to the development of patient-centered guidelines. Foreground questions will be developed based on (i) the hypotheses and conclusions in MASK-air studies and (ii) the search queries on Google Trends.
- (4) AI-assisted questions: LLMs, namely ChatGPT (version 4.0; OpenAI, San Francisco, Calif), will be used for the generation of direct guideline questions (Sousa-Pinto et al.¹⁴⁶). In addition, ChatGPT will be used to help with processing and classifying Google Trends queries. In detail, we will use ChatGPT in 2 different ways for question generation: (i) we will prompt ChatGPT to either assume the role of a patient or of a healthcare provider and provide relevant guideline questions in the PICO format; (ii) we will retrieve popular queries on AR using Google Trends and use ChatGPT to classify these queries into those conveying potentially relevant questions versus those not conveying questions (queries identified as potentially conveying relevant questions will then be manually transformed into guideline questions in the PICO format).

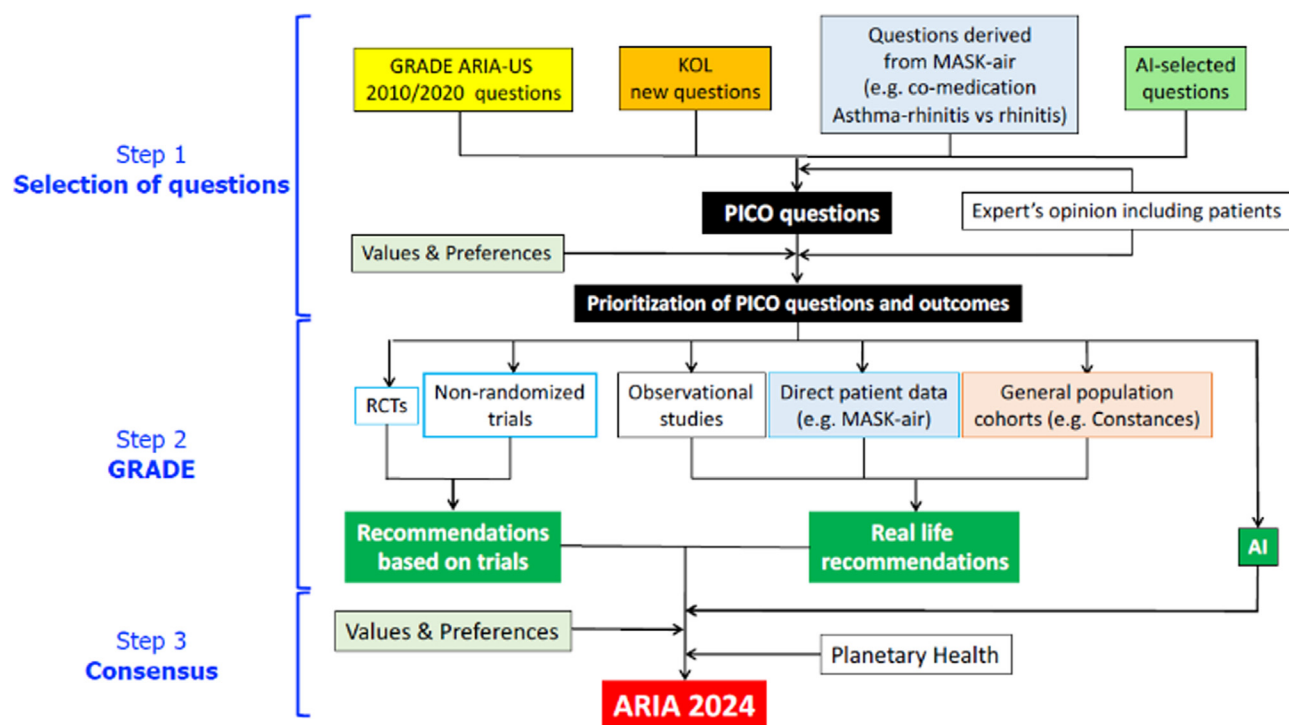


FIGURE 5. Stepwise approach for the development of the ARIA 2024 recommendations. *AI*, Artificial intelligence; *ARIA*, Allergic Rhinitis and its Impact on Asthma; *GRADE*: Grading of Recommendations, Assessment, Development and Evaluation; *KOL*, key opinion leader; *MASK*, Mobile Airways Sentinel network; *PICO*, Population, Interventions, Comparators, and Outcomes; *RCT*, randomized controlled trials.

A consensus meeting to review the set of proposed questions will be held before question prioritization.

Question prioritization. Panel members will be asked to use a visual analog scale (VAS) to rate the priority of each question on a scale of 1 to 9. The ratings will be reviewed by panel co-chairs and the results discussed in a panel meeting. A consensus will then be reached with regard to the questions to be approached in the guidelines.

Outcome generation and prioritization. The questions described previously will also include a list of potential patient-important outcomes identified by the co-chairs, panel suggestions, and a systematic review of patients' values and preferences (Brozek et al.,⁸⁷). Panel members will be asked to use a VAS to rate the priority of each outcome. To ensure that panel members envision the same outcome when discussing the evidence, we will develop health-outcome descriptors to create common definitions that describe the outcomes with respect to symptoms, time horizon, testing and treatment, and consequences (R.J. Vieira, MD, personal communication, 2024).

From evidence to recommendations and digitally enabled, ARIA person-centered care (steps 2 and 3)

For each of the prioritized guideline questions, new or updated systematic reviews of RCTs will be conducted to obtain the best available evidence. In the GRADE approach, incorporation of the best available evidence for the formulation of recommendations involves the use of the EtD framework. The EtD comprises 12 criteria (including, among others, the priority of the problem,

benefits and harms, patients' values and preferences, resource use and cost-effectiveness, impact on health equity, feasibility, and acceptability), enabling each prioritized question to be answered by the formulation of recommendations. When relevant, evidence from RCTs will be complemented by:

- Evidence obtained by observational studies, including the Constances general population cohort or national health data.
- Real-life direct patient data from MASK-air: The database (currently 600,000 days) will be used to provide complementary evidence to the guideline questions, especially for subgroup analyses (eg, considering patients with AR+asthma and AR without asthma), resource use, feasibility, and acceptability of interventions. This aims to incorporate evidence more aligned with the actual experiences of patients in the decision-making process.

Importantly, the EtD framework includes patients' values and preferences as one of the criteria for decision-making. Therefore, we conducted a systematic review to synthesize and appraise all available evidence on patients' values and preferences for health outcomes associated with AR (Brozek et al., submitted), thereby allowing for panel members to issue recommendations aligned with patients' values and preferences.

Consensus to develop the final ARIA 2024 recommendation

A consensus will be made using recommendations obtained from EtDs of RCTs and real-life data (step 3). Importantly, the

EtD framework includes patients' values and preferences as one of the criteria for decision-making. Therefore, we conducted a systematic review to synthesize and appraise all available evidence on patients' values and preferences for health outcomes associated with AR,⁸⁷ thereby allowing for panel members to issue recommendations that are aligned with patients' values and preferences.

ARIA covers all age groups (steps 2b and 3b)

As already done in ARIA 2010, special attention will be paid to children and old-age people, even though the number of RCTs or RWE studies is relatively low. A special subgroup will assess this important topic.

THE POLITICAL AGENDA

Current digital health tools such as MASK-air were developed initially for rhinitis and asthma. However, the technology itself is generic and can be applied to other diseases (eg, the Chronic Urticaria Self Evaluation [CRUSE] mobile app in urticaria).¹⁴⁷ The digital tool enables patients to be guided for an ICP to adapt the medication based on symptom load but also to allow better SDM. The success strongly depends on the patients' adherence, particularly among the elderly, to these digital tools.

Building an alliance among patients, healthcare providers, and policymakers is therefore essential for saving healthcare costs and providing better care for the patients. Healthcare providers or insurers could offer a financial reward to encourage patients with chronic disease to use digital health tools. Allergic diseases are the most frequent chronic diseases in the younger population of industrialized countries. It has been shown that up to €100 billion can be saved every year in socioeconomic costs, mainly due to presenteeism, if patients are correctly treated.² Saving socioeconomic costs will not only present a short-term benefit for the healthcare system but could also result in a strong benefit for society. Therefore, an urgent need also exists for support from policymakers to optimize patient care.

Several policy-focused initiatives have been developed in collaboration with ARIA. They include the Polish Presidency of the EU (2012: Prevention and control of childhood asthma and allergy in the EU from the public health point of view),¹⁴⁸ the Vilnius Declaration (2019: Vilnius Declaration on chronic respiratory diseases: multisectoral care pathways embedding guided self-management, mHealth and air pollution in chronic respiratory diseases),¹⁴⁹ the Finnish Presidency of the EU (2019: Europe that protects),^{52,109,111} and UCRAID (2023: Ukrainian Citizen and refugee electronic support in Respiratory diseases, Allergy, Immunology and Dermatology).¹⁵⁰ Moreover, MASK-air is a best practice of OECD for public health on integrated care for chronic diseases,¹⁵¹ and it has been endorsed by the Ministries of Health of Ukraine (2023)¹⁵⁰ and Poland (2024).

MASK-air has recently been listed as 1 of the 13 OECD best practices of an integrated care model of key strategic interest to policymakers.¹⁵² UCRAID, developed by ARIA and UCARE (Urticaria Centers of Reference and Excellence), is under the auspices of the Ukraine Ministry of Health as well as EAACI, the European Respiratory Society, the European Society of Dermatologic Research, and national societies.¹⁵⁰

A special effort needs to be undertaken to globalize the care pathways. The first ARIA report involved low- and middle-income countries.¹⁵³ A specific group of members of developing countries will be involved in ARIA 2024. Smartphone ownership is growing

rapidly around the world. The International Telecommunication Union (ITU) estimates that approximately 78% of the world's population owns a smartphone, ranging from 47% in low-income countries to 70% in lower-middle-income countries and 94% in high-income countries.¹⁵⁴ The joint WHO-ITU (International Telecommunication Union) initiative "Be He@lthy, Be Mobile" for the prevention and management of noncommunicable diseases, their comorbidities, and their risk factors, including improving disease diagnosis and tracking, is of significant importance. MASK-air is one of the examples of the *Be He@lthy, Be Mobile* handbook on how to implement mBreatheFreely for asthma and chronic obstructive pulmonary disease.¹⁵⁵ The ultimate goal of the initiative would be to propose a "Universal Health Coverage," although this may be beyond the scope of ARIA 2024 (<https://www.who.int/westernpacific/health-topics/detail/universal-health-coverage>).

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ONLINE REPOSITORY

TABLE E1. ARIA strategic overview

Acronym and reference	Name	Dates
WHO-associated projects		
ARIA ^{E1-E5}	Allergic Rhinitis and its Impact on Asthma	1999-
WHO Collaborating Center for Asthma and Rhinitis (Montpellier)		2004-14
GARD ^{E6}	Global Alliance against chronic Respiratory Diseases, demonstration project	2003-23
WHO-ITU ^{E7}	<i>Be He@lthy, Be Mobile</i> handbook on asthma and COPD	2017
EU grants and projects		
GA ² LEN ^{E8}	Global Allergy and Asthma European Network (FP6)	2004-
MeDALL ^{E9,E10}	Mechanisms of the Development of Allergy (FP7)	2009-14
EIP on AHA ^{E11}	European Innovation Partnership on Active and Healthy Ageing (DG Santé and CONNECT)	2012-20
Joint Research Center (JRC) Scientific and Policy Reports on Strategic Intelligence Monitor on Personal Health Systems Phase 3 (SIMP3S) ^{E12}		2015
MACVIA ^{E13}	European Regional Development Fund (ERDF-Région Languedoc-Roussillon)	2016-17
Twinning ^{E14}	Transfer of Innovation (DG Santé and CONNECT)	2017-19
DHE Twinning ^{E15}	Transfer of innovation in severe asthma (H2020)	2019-20
POLLAR ^{E16,E17}	Impact of air Pollution on Asthma and Rhinitis (EIT Health)	2018-19
CATALYSE ^{E18}	Climate change (Horizon Europe)	2022-
MASK@PACA	European Regional Development Fund (ERDF-Région PACA)	2021-22
Good Practice DG Santé on digital health (DG Santé) ^{E19}		2018
Best Practice OECD-DG Santé ^{E20}		2023
ARIA-EAACI Task Forces and projects		
Combined symptom-medication scores for allergic rhinitis (CSMS) ^{E21}		2021
Digital biomarkers in rhinitis and asthma including electronic daily symptom-control score in asthma (e-DASTHMA) ^{E22,E23}		2022
Digitally-enabled, person-centred care (PCC) in allergen immunotherapy: An ARIA-EAACI Position Paper ^{E24}		2023
UCRAID (Ukrainian Citizen and refugee electronic support in Respiratory diseases, Allergy, Immunology and Dermatology) ^{E25}		2023
INTERAID (International Travel Electronic support in Respiratory, Allergy, Immunology and Dermatology)		2024

Updated from Bousquet et al.^{E26}

COPD, Chronic obstructive pulmonary disease; *DG*, Directorate General; *EAAACI*, European Academy of Allergy and Clinical Immunology; *EIT*, European Institute of Innovation and Technology; *OECD-DG*, Organization for Economic Co-operation and Development—Directorate General; *PACA*, Provence-Côte d'Azur; *WHO-ITU*, World Health Organization—International Telecommunication Union.

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