

Alma Mater Studiorum Università di Bologna
Archivio istituzionale della ricerca

Results of an Expert Consensus Survey on the Treatment of Pulmonary Arterial Hypertension With Oral Prostacyclin Pathway Agents

This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Results of an Expert Consensus Survey on the Treatment of Pulmonary Arterial Hypertension With Oral Prostacyclin Pathway Agents / McLaughlin V.V.; Channick R.; De Marco T.; Farber H.W.; Gaine S.; Galie N.; Krasuski R.A.; Preston I.; Souza R.; Coghlan J.G.; Frantz R.P.; Hemnes A.; Kim N.H.; Lang I.M.; Langleben D.; Li M.; Sitbon O.; Tapson V.; Frost A.. - In: CHEST. - ISSN 0012-3692. - ELETTRONICO. - 157:4(2020), pp. 955-965. [10.1016/j.chest.2019.10.043]

Availability:

This version is available at: <https://hdl.handle.net/11585/731907> since: 2020-02-24

Published:

DOI: <http://doi.org/10.1016/j.chest.2019.10.043>

Terms of use:

Some rights reserved. The terms and conditions for the reuse of this version of the manuscript are specified in the publishing policy. For all terms of use and more information see the publisher's website.

This item was downloaded from IRIS Università di Bologna (<https://cris.unibo.it/>).
When citing, please refer to the published version.

(Article begins on next page)

Results of an Expert Consensus Survey on the Treatment of Pulmonary Arterial Hypertension With Oral Prostacyclin Pathway Agents

Vallerie V. McLaughlin, MD, FCCP; Richard Channick, MD; Teresa De Marco, MD; Harrison W. Farber, MD, FCCP; Sean Gaine, MD, PhD; Nazzareno Galié, MD; Richard A. Krasuski, MD; Ioana Preston, MD; Rogerio Souza, MD, PhD; J. Gerry Coghlan, MD; Robert P. Frantz, MD; Anna Hemnes, MD; Nick H. Kim, MD; Irene M. Lang, MD; David Langleben, MD; Mengtao Li, MD; Olivier Sitbon, MD, PhD; Victor Tapson, MD; and Adaani Frost, MD

BACKGROUND: Treatment of pulmonary arterial hypertension (PAH) has evolved substantially over the past two decades and varies according to etiology, functional class (FC), hemodynamic parameters, and other clinical factors. Current guidelines do not provide definitive recommendations regarding the use of oral prostacyclin pathway agents (PPAs) in PAH. To provide guidance on the use of these agents, an expert panel was convened to develop consensus statements for the initiation of oral PPAs in adults with PAH.

METHODS: A systematic literature search was conducted using MEDLINE. The established RAND/University of California Los Angeles appropriateness method, which incorporates the Delphi method and the nominal group technique, was used to create consensus statements. Idiopathic, heritable, repaired congenital heart defect, and drug- or toxin-induced PAH (IPAH+) was considered as one etiologic grouping. The process was focused on the use of oral treprostinil or selexipag in patients with IPAH+ or connective tissue disease-associated PAH and FC II or III symptoms receiving background dual endothelin receptor antagonist/phosphodiesterase type 5 inhibitor therapy.

RESULTS: The panel developed 14 consensus statements regarding the appropriate use of oral PPAs in the target population. The panel identified 13 clinical scenarios in which selexipag may be considered as a treatment option.

CONCLUSIONS: The paucity of clinical evidence overall, and particularly from randomized trials in this setting, creates a gap in knowledge. These consensus statements are intended to aid physicians in navigating treatment options and using oral PPAs in the most appropriate manner in patients with PAH.

CHEST 2019; ■(■):■-■

KEY WORDS: oral prostacyclin; oral treprostinil; prostacyclin pathway agent; pulmonary arterial hypertension; selexipag

ABBREVIATIONS: 6MWD = 6-min walk distance; BNP = brain natriuretic peptide; CHEST = American College of Chest Physicians; CTD = connective tissue disease; ERA = endothelin receptor antagonist; FC = functional class; IPAH+ = idiopathic, heritable, repaired congenital heart defect, and drug- or toxin-induced PAH; NT-proBNP = N-terminal prohormone of BNP; PAH = pulmonary arterial hypertension; PDE5i = phosphodiesterase type 5 inhibitor; PPA = prostacyclin pathway agent; RV = right ventricular; UCLA = University of California Los Angeles

AFFILIATIONS: From the Division of Cardiovascular Medicine (Dr McLaughlin), University of Michigan, Ann Arbor, MI; the Department

of Medicine (Dr Channick), University of California Los Angeles Medical Center, Los Angeles, CA; the Department of Cardiology (Dr De Marco), University of California, San Francisco, CA; the Division of Pulmonary, Critical Care and Sleep Medicine (Drs Farber and Preston), Tufts Medical Center, Boston, MA; The Mater Hospital (Dr Gaine), Dublin, Ireland; the Department of Experimental, Diagnostic and Specialty Medicine-DIMES (Dr Galié), University of Bologna, Bologna, Italy; the Duke University Hospital (Dr Krasuski), Durham, NC; the Pulmonary Department (Dr Souza), Heart Institute, University of São Paulo Medical School, São Paulo, Brazil; the Department of Cardiology (Dr Coghlan), Royal Free Hospital, London,

Currently approved therapies for pulmonary arterial hypertension (PAH) consist of endothelin receptor antagonists (ERAs), phosphodiesterase type 5 inhibitors (PDE5is), a soluble guanylate cyclase stimulator, and oral, inhaled, and parenteral prostacyclin pathway agents (PPAs). Three oral PPAs, treprostinil and beraprost (prostacyclin analogs), and selexipag (a selective prostaglandin I2 receptor agonist), have been developed for the treatment of PAH. Use of beraprost is currently restricted to select Asian countries, and oral treprostinil is not widely available outside North America.

For adult patients with PAH and functional class (FC) II or III symptoms, guidelines recommend initial combination therapy with an ERA and PDE5i.^{1,2} If a patient's risk status is intermediate while receiving combination therapy, the European Society of Cardiology/European Respiratory Society guidelines and the 6th World Symposium on Pulmonary Hypertension proceedings recommend escalation to triple therapy by adding an oral or parenteral PPA.^{1,3} However, discordance exists between the European Society of Cardiology/European Respiratory Society guidelines³ and the American College of Chest Physicians (CHEST) guidelines.² Because the primary end point in the phase 3 oral PPA trials differed, CHEST chose to base their

guidance on 6-min walk distance (6MWD) because it was a common end point among the trials. Based on these data, CHEST² was not able to make a definitive recommendation on when to add selexipag and found no evidence to support the addition of oral treprostinil to ERA and/or PDE5i therapy.

Five randomized trials of oral treprostinil or selexipag have been published.⁴⁻⁸ Two phase 3, placebo-controlled clinical trials (FREEDOM-C and FREEDOM-C2) evaluated oral treprostinil in adults with primarily FC II (23%) or FC III (74%) symptoms receiving combination therapy with an ERA and/or a PDE5i.^{5,7} In contrast with the phase 3 study of oral treprostinil monotherapy in which patients treated with oral treprostinil vs placebo demonstrated a statistically significant improvement in 6MWD,⁶ no statistically significant improvement in the primary end point (6MWD) was observed in the combination therapy trials.^{5,7} In a phase 2 trial comparing the addition of selexipag vs placebo in adults receiving combination therapy with an ERA and/or a PDE5i, patients treated with selexipag had a statistically significant decrease in mean pulmonary vascular resistance, the primary end point of the study.⁴ One phase 3 randomized, placebo-controlled clinical trial (GRIPHON) evaluated selexipag in adults with primarily FC II (46%) or FC III (53%) symptoms receiving background therapy with an ERA and/or PDE5i or no background therapy.⁸ The primary end point was a composite of death or a complication related to PAH, which included disease progression or worsening of PAH that resulted in hospitalization, initiation of parenteral PPA therapy or long-term oxygen therapy, or the need for balloon atrial septostomy or lung transplantation. In the overall patient population, selexipag statistically significantly reduced the risk of a primary end point event. In subgroup analysis, the treatment effect was consistent among subgroups regardless of background therapy.

The gaps in data with oral PPAs in patients with FC II or III symptoms who are receiving ERA plus PDE5i coupled with a lack of definitive guidance on their use in published treatment guidelines creates uncertainty for physicians regarding the role of oral PPAs in managing these patients. Additionally, complicated real-world scenarios arise that are not well addressed in the setting of a controlled clinical trial. Therefore, we sought to use a well-described scientific methodology to develop expert consensus opinion statements on when to initiate the oral PPAs treprostinil and selexipag in common

England; the Department of Cardiovascular Medicine (Dr Frantz), Mayo Clinic College of Medicine and Science, Rochester, MN; the Division of Allergy, Pulmonary and Critical Care Medicine (Dr Hemnes), Vanderbilt University Medical Center, Nashville, TN; the Department of Cardiothoracic Surgery (Dr Kim), University of California San Diego Medical Center, San Diego, CA; the Department of Internal Medicine II (Dr Lang), Division of Cardiology, Medical University of Vienna, Allgemeines Krankenhaus, Vienna, Austria; the Center for Pulmonary Vascular Disease (Dr Langleben), Cardiology Division, Jewish General Hospital and McGill University, Montreal, QC, Canada; the Department of Rheumatology (Dr Li), Peking Union Medical College Hospital, Peking Union Medical College, Chinese Academy of Medical Sciences, Beijing, China; the Université Paris-Saclay (Dr Sitbon), APHP, Hôpital Bicêtre, Service de Pneumologie, Le Kremlin-Bicêtre, France; the Department of Medicine (Dr Tapson), Cedars-Sinai Medical Center, Los Angeles, CA; and the Department of Medicine (Dr Frost), Institute of Academic Medicine, Houston Methodist Hospital, Houston, TX.

Dr Frost is now retired.

This work has been presented as a late-breaking poster at the CHEST Annual Meeting, October 19-23, 2019, New Orleans, LA.

FUNDING/SUPPORT: Actelion Pharmaceuticals provided funding that supported use of independent providers of Delphi methodology expertise and nominal group technique, survey creation, data analysis, medical communication, and meeting management.

CORRESPONDENCE TO: Vallerie V. McLaughlin, MD, FCCP, Pulmonary Hypertension Program, University of Michigan Cardiovascular Center, 1500 E Medical Center Dr, SPC 5853, Ann Arbor, MI 48109; e-mail: vmclaugh@umich.edu

Copyright © 2019 American College of Chest Physicians. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

DOI: <https://doi.org/10.1016/j.chest.2019.10.043>

clinical scenarios in adults with PAH and World Health Organization FC II or III symptoms.

Expert consensus statements cannot replace assessment and clinical decision-making by a qualified health-care practitioner for an individual patient. These statements are intended to guide physicians in common scenarios and do not address all possible clinical situations, nor do these statements account for

additional individual patient factors not specifically stated, such as various comorbidities, patient preference, the ability of a patient to manage or adhere to a treatment, or the patient's ability to pay for treatment. Additionally, the consensus statements presented within are not intended for use as criteria for third-party payor reimbursement of specific drugs or treatments for groups or individuals with PAH of any etiology.

Methods

The consensus process is outlined in [Figure 1](#). The panel used the established RAND/University of California Los Angeles (UCLA) appropriateness method,⁹ which incorporates the Delphi method and the nominal group technique, to create consensus statements. This method was developed to reach consensus among participants, particularly in situations in which evidence is lacking to support decision-making. The process was directed by a moderator skilled in the RAND/UCLA appropriateness method. All authors served as panel members and were chosen by a small group of PAH experts who selected members with a goal of representing a variety of geographic regions and clinical expertise both in patients with PAH and in the use of oral PPAs. Data were analyzed independently by Humanitas, Inc (Silver Spring, MD).

Funding was provided by Actelion Pharmaceuticals. This funding supported the use of independent providers of Delphi methodology expertise and nominal group technique, survey creation, data analysis, medical communication, and meeting management. The authors were not paid honoraria for their participation. Actelion Pharmaceuticals played no role in the literature search and analysis, development of surveys used to gather consensus, or data analysis; and no Actelion Pharmaceuticals employee was present at the face-to-face meeting during which consensus statements were finalized. The current paper was drafted, critically reviewed, and edited solely

by the authors with support from an independent professional medical communications agency. Actelion Pharmaceuticals reviewed the final manuscript only to ensure accuracy of selexipag background information; no edits were made to the manuscript based on this review.

A systematic literature search was conducted using MEDLINE via PubMed using the following search terms: (“pulmonary arterial hypertension” OR “pulmonary hypertension”) AND (prostacyclin [tw] OR prostanoid[tw] OR PGI2[tw]). The search was limited to English language, adult patients (≥ 18 years of age), group 1 PH (ie, PAH), human clinical studies, and a 10-year time frame from October 1, 2008, to October 1, 2018. Relevant articles containing clinical information and review articles were retained. The search was augmented with drug prescribing information for PPAs (epoprostenol injection; treprostinil tablets, inhalation, and injection; iloprost inhalation; and selexipag tablets), key articles identified in reference lists outside the search time window, and pivotal trials for oral treprostinil and selexipag (additional details are provided in [e-Appendix 1](#)).

Based on the literature search, and informed by expert opinion and an initial presurvey of the panelists, the first author and the moderator decided that questions would be posed about oral treprostinil and selexipag separately, that idiopathic, heritable, repaired congenital heart defect, and drug- or toxin-induced PAH (IPAH+) would be

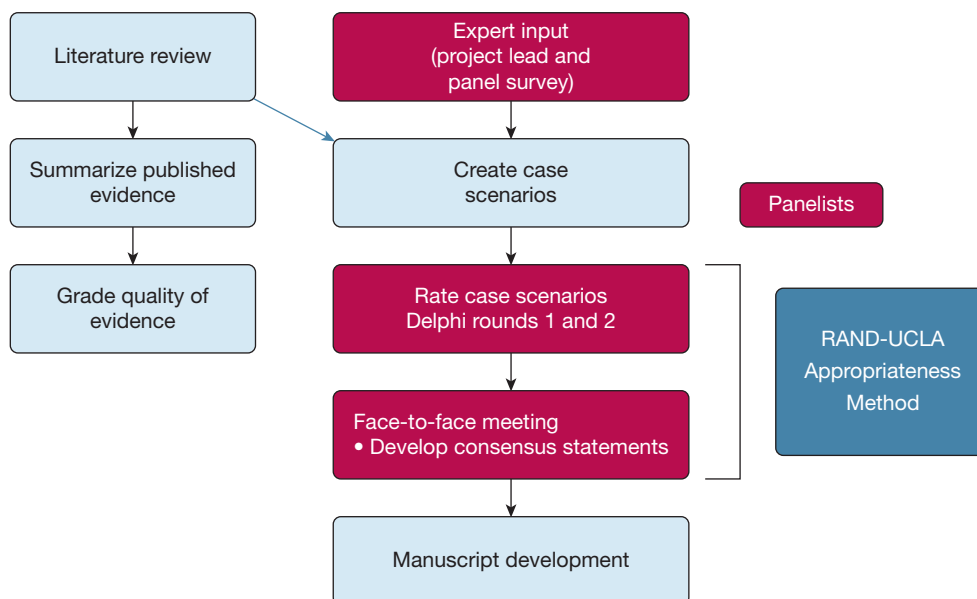


Figure 1 – Overview of the consensus methodology. UCLA = University of California Los Angeles.

TABLE 1] Hemodynamic Values Comprising Low-, Intermediate-, and High-Risk Groups

Hemodynamic Parameter	Low	Intermediate	High
Right atrial pressure, mm Hg	< 8	8-14	> 14
Cardiac index, L/min/m ²	≥ 2.5	2.0-2.4	< 2.0
Mixed venous oxygen saturation, %	> 65	60-65	< 60

considered as one etiologic grouping, that the process would focus on the use of oral treprostinil or selexipag in patients on background dual ERA/PDE5i therapy with FC II or III symptoms, and that the process would exclude oral PPA monotherapy, upfront double combination therapy with an oral PPA and another agent, and use of an oral PPA in patients with FC IV PAH, consistent with current evidence and clinical practice. Panelists ranked, in descending order of importance, the clinical factors that they typically use to make routine treatment decisions regarding the initiation of oral PPAs. The initial list of clinical factors was drawn from clinical trial end points and multiparameter risk assessment algorithms. Based on these results, the following clinical factors were considered in this order of importance (within each FC): hemodynamics, PAH-associated hospitalization within the prior 6 months, right ventricular (RV) function, serum brain natriuretic peptide (BNP)/N-terminal prohormone of BNP (NT-proBNP) levels, and 6MWD. Although PAH-associated hospitalization was ranked higher than hemodynamics in the survey, the first author and the moderator opted to construct the survey so that the panel would consider hemodynamics first, reasoning that these data are more likely to be available to the physician at the time of decision-making and because hemodynamics have historically been the most critical factor in decision-making. Clinical factors evaluated but excluded from further ranking were right atrial area, stroke volume index, age, sex, cardiopulmonary exercise testing, diffusing capacity of the lungs for carbon monoxide, BP, heart rate, clinically significant renal insufficiency, syncope, and RV failure (the last two are indicative of FC IV, where evidence supports the use of parenteral prostacyclin therapy^{2,3}).

Panelists were presented with a series of clinical scenarios created by the first author and the moderator for a patient in one of three etiologic groups (IPAH+, connective tissue disease [CTD]-associated PAH, and portopulmonary hypertension), with FC II or III symptoms, and with mostly low-, intermediate-, or high-risk hemodynamic parameters (based on two of the three following variables meeting the risk category level: right atrial pressure, mixed venous oxygen saturation, and cardiac index) (Table 1). Panelists were then asked questions sequentially about the appropriateness of selexipag or oral treprostinil in patients with a specific clinical scenario regarding clinical factors in the following order: (1) hospitalization because of PAH in the last 6 months (yes or no), (2) RV function (normal, mild dysfunction, or moderate/severe dysfunction based on echocardiogram or MRI), (3) BNP/NT-proBNP levels (normal or abnormal), and (4) 6MWD (> 440 or ≤

440 m) (e-Fig 1). Consistent with the RAND/UCLA method, cost was not considered in the decision-making model.

For Delphi round 1 (Delphi 1), 1,620 case scenarios were presented with an equal number for treprostinil and selexipag, respectively. Panelists assigned a score of 1 to 9 for each scenario, with scores of 1 to 3 indicating that the oral PPA therapy is inappropriate for that patient scenario with risks outweighing benefits, scores of 4 to 6 indicating the risk to benefit ratio is uncertain and decisions are made on an individual basis, and scores of 7 to 9 indicating the therapy is appropriate and benefits clearly outweigh the risks. If a respondent assigned a score of 1 to 3 for a scenario, the software cut off further downstream questions and a score of 2 (ie, mean and median of 1 to 3) was imputed for that individual participant. Mean, median, mode, and response distribution according to bottom, middle, and top third of the scale (1 to 3, 4 to 6, and 7 to 9) were calculated. For a case scenario to be included for reassessment in Delphi 2, a threshold median score of > 3 had to be met. Case scenarios with a median score ≤ 3 were interpreted as a consensus against the appropriateness of the oral PPA in that case. A total of 677 case scenarios passed Delphi 1. For included case scenarios, data for individual respondents were gathered in a summary along with their individual score for each scenario, the median score, and the frequency distribution. This summary was sent to the panelists with the Delphi 2 survey. Panelists had the opportunity to retain their original response or to change their response after seeing the group median and score distribution for each question. In Delphi 2, the threshold for preliminary consensus agreement for appropriateness of the oral PPA in a case scenario was a median score ≥ 7 with ≤ 33% of respondents scoring the appropriateness as 1 to 3. Items with a median score < 7 were designated as case scenarios lacking consensus agreement (and therefore rejected). Items with median score ≥ 7 and with > 33% scored 1 to 3 were discussed at the face-to-face meeting, during which nominal group technique was used to obtain group consensus on each of the draft consensus statements developed based on Delphi 2. Each consensus statement was discussed and voted on silently using a computerized audience response system and the same scale (1 to 9, with 1 indicating not appropriate). Panelists agreed to not designate the strength of evidence for the consensus statements given the paucity of clinical and/or trial evidence for the clinical scenarios analyzed. The RAND/UCLA method is designed to gain consensus in situations with insufficient evidence, such as clinical scenarios not well represented in clinical trials.

Results

A total of 677 case scenarios passed Delphi 1, and 458 were accepted in Delphi 2. During discussion, the panel determined that the final clinical factor rankings were most appropriate for patients with IPAH+ and CTD-associated PAH and drafted 14 consensus statements (Table 2). Consensus statements for use of oral PPAs in portopulmonary hypertension were not developed

because treatment goals were ambiguous: symptom improvement vs achievement of transplant-acceptable hemodynamic thresholds.

The median score for the use of oral treprostinil did not meet the predetermined threshold for a recommendation in favor of its use in any clinical scenario evaluated (median scores all < 7). Oral

TABLE 2] Expert Consensus Statements

Patients With IPAH+ and Low- or Intermediate-Risk Hemodynamic Parameters	
Selexipag may be considered for patients with IPAH+ who are receiving dual oral therapy with an ERA and PDE5i and who meet any of the following clinical scenario criteria:	
FC II	
1	FC II symptoms and low-risk hemodynamic parameters, and who <i>have not</i> been hospitalized for PAH in the last 6 mo, but have moderate-to-severe RV dysfunction, irrespective of their BNP/NT-proBNP levels, or 6MWD
2	FC II symptoms and low-risk hemodynamic parameters, and who <i>have</i> been hospitalized for PAH in the last 6 mo, irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD
3	FC II symptoms and intermediate-risk hemodynamic parameters, irrespective of hospitalization for PAH in the last 6 mo, their RV function, BNP/NT-proBNP levels, or 6MWD
FC III	
4	FC III symptoms and low-risk hemodynamic parameters irrespective of hospitalization for PAH in the last 6 mo, their RV function, BNP/NT-proBNP levels, or 6MWD
5	FC III symptoms and intermediate-risk hemodynamic parameters, who <i>have not</i> been hospitalized for PAH in the last 6 mo, and irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD
6	FC III symptoms and intermediate-risk hemodynamic parameters who <i>have</i> been hospitalized for PAH in the last 6 mo, and with normal or mildly impaired RV function, irrespective of BNP/NT-proBNP levels, or 6MWD
Patients With CTD-Associated PAH and Low- or Intermediate-Risk Hemodynamic Parameters	
Selexipag may be considered for patients with CTD-associated PAH and one of the following clinical scenarios:	
FC II	
7	FC II symptoms and low-risk hemodynamic parameters, and who <i>have not</i> been hospitalized for PAH in the last 6 mo, but have any degree of RV dysfunction and abnormal BNP/NT-proBNP levels, irrespective of 6MWD
8	FC II symptoms and low-risk hemodynamic parameters, and who <i>have</i> been hospitalized for PAH in the last 6 mo irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD
9	FC II symptoms and intermediate-risk hemodynamic parameters, irrespective of hospitalization for PAH in the last 6 mo, their RV function, BNP/NT-proBNP levels, or 6MWD
FC III	
10	FC III symptoms and low-risk hemodynamic parameters, and who <i>have not</i> been hospitalized for PAH in the last 6 mo, and RV function is abnormal, BNP/NT-proBNP levels are abnormal, or 6MWD is ≤ 440 m
11	FC III symptoms and low-risk hemodynamic parameters, and who <i>have</i> been hospitalized for PAH in the last 6 mo, irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD
12	FC III symptoms and intermediate-risk hemodynamic parameters, who <i>have not</i> been hospitalized for PAH in the last 6 mo, and irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD
13	FC III symptoms and intermediate-risk hemodynamic parameters who <i>have</i> been hospitalized for PAH in the last 6 mo, and with normal or mildly impaired RV function, irrespective of BNP/NT-proBNP levels, or 6MWD
Patients With IPAH+ or CTD-Associated PAH and High-Risk Hemodynamic Parameters	
14	In patients with idiopathic, heritable, drug- or toxin-induced, or repaired congenital heart disease-associated PAH or connective tissue disease-associated PAH who are on dual oral ERA/PDE5i therapy and who have high-risk hemodynamic parameters, IV or subcutaneous prostacyclin is the treatment of choice

6MWD = 6-min walk distance; BNP = B-type natriuretic peptide; CTD = connective tissue disease; ERA = endothelin receptor antagonist; FC = World Health Organization functional class; IPAH+ = idiopathic, heritable, drug- or toxin-induced, or repaired congenital heart disease-associated PAH; NT-proBNP = N-terminal prohormone BNP; PAH = pulmonary arterial hypertension; PDE5i = phosphodiesterase type 5 inhibitor; RV = right ventricular.

treprostinil data were analyzed according to panel members' location (United States vs non-United States) to examine the possibility that the lack of oral treprostinil availability in non-US locations may have affected the appropriateness determination. However, all scenarios that entered Delphi 2 were rejected in both US and non-US panelist subgroups. Median scores for all scenarios ranged from 3 to 6.5 among US panelists and from 1.5 to 2.5 among non-US panelists. Therefore, the

resulting panel consensus statements are limited to the use of clinical situations in which oral selexipag is considered appropriate.

IPAH+ and Low- or Intermediate-Risk Hemodynamics

Among patients with IPAH+ and low- or intermediate-risk hemodynamic parameters who are receiving dual oral ERA/PDE5i therapy, the panel determined that

selexipag may be considered as additional therapy for patients with the following clinical scenarios (Fig 2).

FC II: The panel determined that in patients with IPAH+ and FC II symptoms, selexipag may be considered in patients with low-risk hemodynamics, if the patient has not been hospitalized for PAH in the last 6 months but has moderate-to-severe RV dysfunction, irrespective of their BNP/NT-proBNP levels, or 6MWD. The panelists discussed that severe RV dysfunction may represent a poor prognostic factor¹⁰⁻¹³ and determined that RV dysfunction in this patient population warrants additional therapy with selexipag.

In patients with IPAH+ and FC II symptoms, selexipag may be considered in patients with low-risk hemodynamics if the patient has been hospitalized for PAH in the last 6 months, irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD. The panelists discussed that patients in clinical practice who fit this scenario may be younger or those with drug- or toxin-induced PAH whose noncompliance with diuretics have led to hospitalization. The panel agreed that hospitalization for worsening PAH represents a poor prognostic factor,¹⁴ and the patient may require additional medication.

In patients with IPAH+ and FC II symptoms, selexipag may be considered in patients with intermediate-risk hemodynamics, irrespective of hospitalization for PAH in the last 6 months, RV function, BNP/NT-proBNP levels, or 6MWD.

FC III: The panel determined that in patients with IPAH+ and FC III symptoms, selexipag may be considered in patients with low-risk hemodynamics irrespective of hospitalization for PAH in the last 6 months, RV function, BNP/NT-proBNP levels, or 6MWD.

In patients with IPAH+ and FC III symptoms, selexipag may be considered in patients with intermediate-risk hemodynamics who have not been hospitalized for PAH in the last 6 months, and irrespective of RV function, BNP/NT-proBNP levels, or 6MWD.

In patients with IPAH+ and FC III symptoms, selexipag may be considered in patients with intermediate-risk hemodynamics who have been hospitalized for PAH in the last 6 months, and with normal or mildly impaired RV function, irrespective of BNP/NT-proBNP levels, or 6MWD. The panelists discussed that this subgroup of hospitalized patients may be appropriate candidates for oral rather than parenteral therapy.

CTD-Associated PAH and Low- or Intermediate-Risk Hemodynamics

Among patients with CTD-associated PAH and low- or intermediate-risk hemodynamic parameters who are receiving dual oral ERA/PDE5i therapy, the panel determined that selexipag may be considered as additional therapy for patients with specific clinical scenarios (Fig 3).

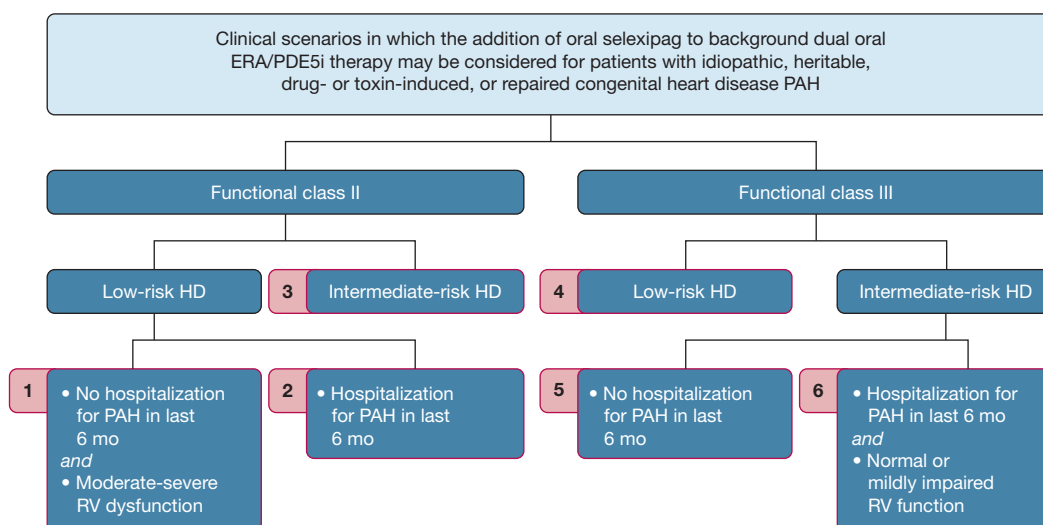


Figure 2 – Clinical scenarios in which the expert panel determined that oral selexipag may be considered for patients with idiopathic, heritable, drug- or toxin-induced, or repaired congenital heart disease-associated PAH who are receiving dual oral therapy with an ERA and a PDE5i. Numbers indicate corresponding consensus statement. ERA = endothelin receptor antagonist; HD = hemodynamic parameters; PAH = pulmonary arterial hypertension; PDE5i = phosphodiesterase type 5 inhibitor; RV = right ventricular.

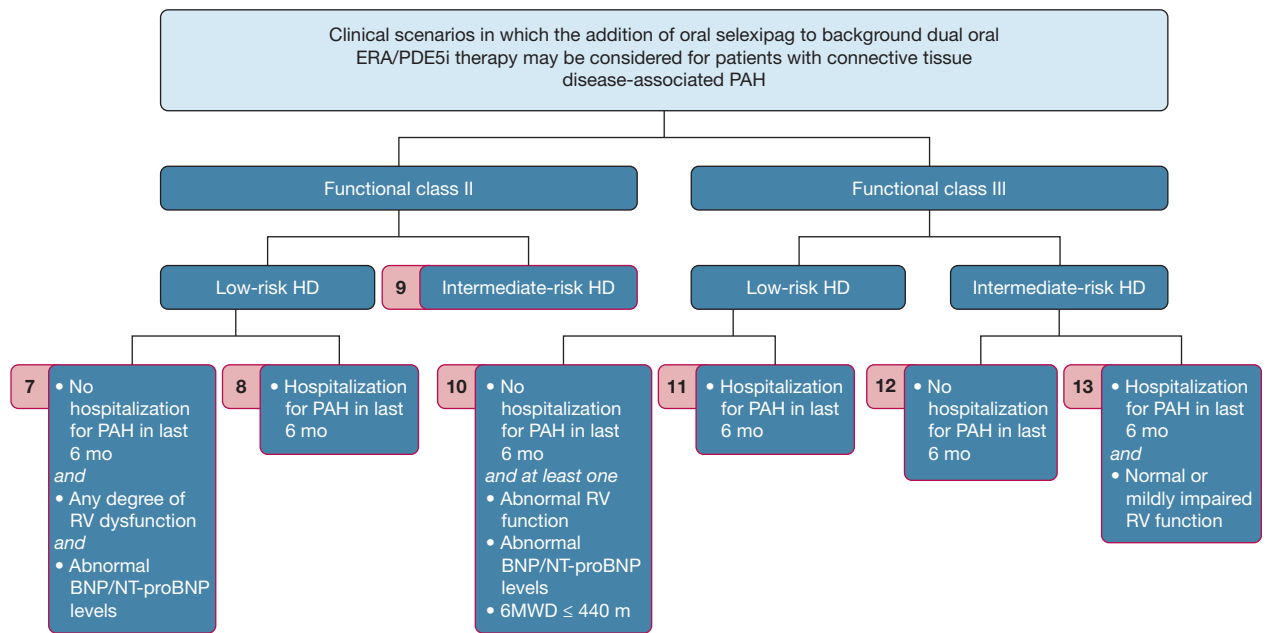


Figure 3 – Clinical scenarios in which the expert panel determined that oral selexipag may be considered for patients with connective tissue disease-associated PAH who are receiving dual oral therapy with an ERA and a PDE5i. Numbers indicate corresponding consensus statement. 6MWD = 6-min walk distance; BNP = B-type natriuretic peptide; NT-proBNP = N-terminal prohormone BNP. See Figure 2 legend for expansion of other abbreviations.

FC II: The panel agreed that in patients with CTD-associated PAH and FC II symptoms, selexipag may be considered for patients with low-risk hemodynamics who have not been hospitalized for PAH in the last 6 months but have any degree of RV dysfunction and abnormal BNP/NT-proBNP levels, irrespective of 6MWD.

In patients with CTD-associated PAH and FC II symptoms, selexipag may be considered for patients with low-risk hemodynamics who have been hospitalized for PAH in the last 6 months, irrespective of RV dysfunction, BNP/NT-proBNP levels, or 6MWD. Panelists noted that some patients in this category may benefit from parenteral therapy, specifically those with moderate-to-severe RV dysfunction and 6MWD \leq 440 m. However, patients with CTD-associated PAH may have difficulty managing parenteral therapy because of the necessity of manipulating pumps and a higher incidence of adverse events compared with patients with IPA⁺¹⁵; selexipag offers an alternative therapy in such situations.

In patients with CTD-associated PAH and FC II symptoms, selexipag may be considered in patients with intermediate-risk hemodynamics, irrespective of hospitalization for PAH in the last 6 months, RV function, BNP/NT-proBNP levels, or 6MWD.

FC III: The panel determined that in patients with CTD-associated PAH and FC III symptoms, selexipag may be considered in patients with low-risk hemodynamics who have not been hospitalized for PAH in the last 6 months, and with abnormal RV function, abnormal BNP/NT-proBNP levels, or 6MWD \leq 440 m.

In patients with CTD-associated PAH and FC III symptoms, selexipag may be considered in patients with low-risk hemodynamics who have been hospitalized for PAH in the last 6 months, irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD.

In patients with CTD-associated PAH and FC III symptoms, selexipag may be considered in patients with intermediate-risk hemodynamics who have not been hospitalized for PAH in the last 6 months, and irrespective of their RV function, BNP/NT-proBNP levels, or 6MWD.

In patients with CTD-associated PAH and FC III symptoms, selexipag may be considered in patients with intermediate-risk hemodynamics who have been hospitalized for PAH in the last 6 months, and with normal or mildly impaired RV function, irrespective of BNP/NT-proBNP levels or 6MWD. Consistent with patients of IPA⁺ etiology, the panelists concluded that this lower-risk subgroup of hospitalized patients may be appropriate candidates for oral rather than parenteral therapy.

IPAH+ or CTD-Associated PAH and High-Risk Hemodynamics

The panel determined that in patients with IPAH+ or CTD-associated PAH, who are on dual oral therapies and who have high-risk hemodynamics, IV or subcutaneous prostacyclin is the treatment of choice. Exceptions may occur in which selexipag may be considered in patients who decline or are unable to use parenteral therapy; each patient must be evaluated individually with consideration for patient preference, comorbidities, and other pertinent patient-related factors.

Discussion

The current expert panel consensus survey represents the first comprehensive set of consensus statements focused on the use of oral PPAs in patients with PAH developed through established, formal methodology for consensus building. These consensus statements were limited to individuals receiving background dual oral therapy with ERA and PDE5i in alignment with current standard of care.¹⁻³ At the time of the literature analysis, data did not support the use of an oral PPA as a component of upfront double combination therapy.

Consistent with treatment guidelines,¹⁻³ the addition of oral PPAs was not recommended for patients with FC IV PAH or high-risk hemodynamics, with parenteral prostacyclin being the treatment of choice for these patients. The addition of selexipag was recommended for patients with FC II IPAH+ and intermediate-risk hemodynamics, in patients with FC III IPAH+ and low-risk hemodynamics, and in patients with FC II CTD-associated PAH and intermediate-risk hemodynamics, irrespective of any other clinical factor, underlining the importance the panel placed on hemodynamic data. Recent hospitalization was also a critical factor for the panel. It was the sole precipitating factor for adding selexipag in patients with FC II IPAH+ or FC II CTD-associated PAH and low-risk hemodynamics, and lack of recent hospitalization was the sole precipitating factor for adding selexipag in patients with FC III IPAH+ or FC III CTD-associated PAH and intermediate-risk hemodynamics. Notably, 6MWD was not a determining factor in recommendations for IPAH+ or CTD-associated PAH, and BNP levels were not considered a key factor in IPAH+.

Some differences emerged in recommendations between treatment of patients with IPAH+ and CTD-associated PAH despite identical clinical scenario queries. First, for

patients with FC II symptoms, low-risk hemodynamics, and no hospitalization in the last 6 months for PAH, experts determined that selexipag could be considered for patients with IPAH+ if they had moderate-to-severe RV dysfunction, but for patients with CTD-associated PAH, it would be considered if they had any degree of RV dysfunction and abnormal BNP/NT-proBNP levels. Second, for patients with FC III symptoms and low-risk hemodynamics, experts determined that selexipag could be considered for patients with IPAH+ irrespective of hospitalization in the last 6 months for PAH and other features, whereas for CTD-associated PAH, patients with no hospitalization in the last 6 months for PAH should have either abnormal RV function, abnormal BNP/NT-proBNP levels, or 6MWD \leq 440 m. A possible explanation for the lower threshold for use in patients with FC II symptoms may be because of the poorer prognosis associated with scleroderma-associated PAH,^{16,17} the panelists perceived a need for more intensive therapy. However, the threshold for addition of selexipag appears to be more nuanced in patients with CTD-PAH vs IPAH+ with FC III symptoms and low-risk hemodynamics. Selexipag use is recommended for patients with IPAH+ in this scenario regardless of hospitalization; however, in patients with CTD, it is recommended if they had been hospitalized or, if that were not the case, if they had another abnormal variable. This scenario suggests a higher threshold for use in patients with CTD-PAH such that another abnormality in a PAH-related variable (in addition to FC III status) is required because there could be an alternate cause of FC III status, such as deconditioning or musculoskeletal limitations, in these patients. A separate consideration among patients with CTD-associated PAH is the frequent presence of GI symptoms, which may affect treatment choice because oral PPAs are known to be associated with GI side effects.⁴⁻⁸ Panelists were not queried about their reasons for voting on scenarios; therefore, explanations for these differences are purely speculative.

Given the availability of a number of different treatment options for PAH, treatment decisions have become more complex. The opinions described should be considered as suggestions, with the caveat that each patient presents a unique set of features that must also be considered. For example, if a patient has not attained treatment goals with the inclusion of an oral PPA, a parenteral PPA will likely be necessary. In addition, there are two inhaled PPAs approved for the treatment of PAH; they may be an alternative to oral PPAs in patients who have

difficulty tolerating them or who prefer an inhaled formulation for other reasons.

Ultimately, these consensus statements were limited to use of oral selexipag because the panel did not support the use of oral treprostinil for the presented clinical scenarios, consistent with published phase 3 clinical trial data at the time of the meeting.^{5,7} Neither FREEDOM-C nor FREEDOM-C2 demonstrated a benefit in the primary end point of 6MWD with the addition of oral treprostinil to background ERA and/or PDE5i. Clinical trial data with oral treprostinil in patients on background monotherapy with either ERA or PDE5i have been reported (FREEDOM-EV¹⁸). When these results are published, they may affect the consensus opinion. However, it should be noted that FREEDOM-EV evaluates the addition of oral treprostinil to monotherapy with an ERA or a PDE5i (ie, double combination therapy). This consensus opinion is built around the addition of an oral PPA to the regimen of combination therapy with an ERA and a PDE5i (ie, triple combination therapy). As such, results of FREEDOM-EV would not have direct bearing on these opinions. Nonetheless, future expert panels should consider these results, along with any additional new data in creating future consensus statements or recommendations.

Patients with portopulmonary hypertension were not considered in the final consensus statements because the goals for treatment differ from those with IPAH+ or CTD-associated PAH. Whether patients are candidates for transplant is an overarching factor in treatment planning, and that line of questioning was not included in the Delphi process. Panelists agree that future consideration of consensus statements for oral PPA use in this population is warranted.

Based on the panel's rankings, clinical factors were considered within each FC. The panel recognizes the subjectivity of this assessment and inherent

disagreement among practicing physicians in assigning FC to an individual patient.¹⁹ For all patients in whom selexipag was considered appropriate, and particularly in those with more high-risk features within an FC category, the panelists made the determination with the expectation of timely and consistent patient follow-up to assess efficacy and to adjust treatment if necessary. The panel also acknowledges that the importance of any given clinical factor is patient-specific and that physicians must use clinical judgment and their knowledge of the individual patient to prioritize clinical factors when making treatment decisions. Data gleaned from this survey highlight the importance of multiparameter risk assessment and its impact on daily clinical decision-making.

A strength of this expert consensus panel survey was the use of the RAND/UCLA method, which is a well-established, reliable, and widely used process for gaining expert consensus in the setting of limited available data. Limitations of the survey include the small quantity of evidence available for evaluation, which was five clinical trials, including two studies of selexipag and three studies of oral treprostinil.⁴⁻⁸ Expert consensus, based on available clinical trial data at the time the survey was conducted, eliminated oral treprostinil as an appropriate therapy for patients with FC II or III IPAH+ or CTD-associated PAH who were receiving an ERA and PDE5i. As such, the resulting consensus statements are based on clinical experience driven by the two primary studies of selexipag.^{4,8}

In summary, this expert panel survey provides physicians with guidance for the use of oral PPAs in patients with FC II or III IPAH+ and CTD-associated PAH receiving dual oral ERA/PDE5i therapy. The paucity of clinical evidence in this setting creates a gap in knowledge. These expert opinions must be validated with rigorous prospective studies, and this document may serve as a template for future investigations.

Acknowledgments

Author contributions: All authors completed the PIXEL surveys, had access to the data, contributed to data interpretation and manuscript writing, provided final approval of the manuscript for submission, and have agreed to be accountable for the work in that any questions concerning the work are investigated and resolved.

Financial/nonfinancial disclosures: The authors have reported to *CHEST* the following: V. V. M. is a consultant and/or advisor for Actelion Pharmaceuticals US, Inc, Bayer Corp, Gilead Sciences, Inc, St. Jude Medical, Steadymed Therapeutics, and United Therapeutics Corp. The University of Michigan has received research funding from Actelion Pharmaceuticals US, Inc, Arena Pharmaceuticals, Bayer Corp, Gilead Sciences, Inc, and Eiger BioPharmaceuticals. R. C. has received researched grants from Actelion Pharmaceuticals US, Inc, and Bayer Corp; and is a consultant to Actelion Pharmaceuticals US, Inc, Bayer Corp, ZappRx, Inc, and ThirdPole. T. D. M. is a consultant for Johnson & Johnson/Actelion Pharmaceuticals, United Therapeutics, Arena, and SCOPE/Bial. She is a speaker for Actelion Pharmaceuticals. H. W. F. has received honoraria from Actelion Pharmaceuticals US, Inc, Bayer Corp, and SAB Biotherapeutics. He is a consultant for Actelion Pharmaceuticals, United Therapeutics, Boehringer-Ingelheim, and Bristol-Myers Squibb. He has performed end point adjudication for United Therapeutics. S. G. has received honoraria from Actelion Pharmaceuticals and United Therapeutics. He has received travel grants from Actelion, Novartis, and Menerini. He has performed drug safety board monitoring for United Therapeutics, GSK, and Novartis. N. G. reports grants and personal fees from Actelion, Bayer, GSK, and Pfizer; and personal fees from MSD, all outside the submitted work. R. A. K. is a consultant and receives research funding from Actelion Pharmaceuticals. He is also an investigator for Edwards Lifesciences and is an unpaid member of the scientific advisory board for Ventripoint. I. P. has been a principle investigator on studies sponsored by Actelion, Acceleron, Bayer, Complexa, Liquidia, PhaseBio, Tenax, and United Therapeutics. She has been a steering committee member for Actelion, Acceleron, and Liquidia; and an adjudication committee member for Pfizer. She has served as a consultant for Actelion, Respira, and United Therapeutics; and has reviewed grants for Gilead and United Therapeutics. R. S. has received consultancy fees from Actelion, Bayer, Acceleron, GSK, and Pfizer. J. G. C. has received consultancy fees and honoraria from Actelion Pharmaceuticals, United Therapeutics, GSK, and Bayer; study grants from Actelion Pharmaceuticals; and conference fees from Bayer. R. P. F. is a consultant and steering committee member for Actelion Pharmaceuticals US, Inc; and has served on advisory boards for Abbott and

United Therapeutics. A. H. has received grants from the NIH and CMREF. She has served as a consultant for Actelion, Bayer, Complexa, United Therapeutics, and PHPrecisionMed where she also owns equity interest. N. H. K. has received research support from Bellerophon, Eiger, Gossamer Bio, Lung Biotechnology, and SoniVie. He has served as a consultant for Actelion, Arena, Bayer, MSD, and United Therapeutics. He has served on the speakers bureaus for Actelion and Bayer. I. M. L. has relationships with pharmaceutical companies including AOPOrphan Pharmaceuticals AG, Actelion-Janssen (speaker honoraria and grants to the institution), MSD (speaker honoraria), Medtronic (speaker honoraria and travel expenses), and Ferrer (speaker honoraria and travel expenses). D. L. is a consultant and on advisory boards for Acceleron, Actelion, Bayer, Bellerophon, Northern Therapeutics, and PhaseBio. He has received research support from Actelion, Bayer, Eiger, Northern Therapeutics, Reata, and United Therapeutics. He has been on speakers bureaus for Actelion and Bayer. He has received travel and accommodation support from Acceleron, Actelion, Bayer, and Northern Therapeutics. O. S. has received research grants from Actelion, Bayer, GlaxoSmithKline, and MSD. He also has served as a consultant to Actelion, Bayer, Ferrer, Gossamer Bio, MSD, and United Therapeutics. V. T. has served on advisory boards and has received research funding from Actelion, Bayer, and United Therapeutics. He has served on the steering committees for the latter companies, and for Vwave. A. F. has received honoraria and travel expenses for Actelion-sponsored lectures and expert advice litigation related to PH therapies; has received honoraria for consultation and study end point adjudication for United Therapeutics; consults for PhaseBio; and performs data safety monitoring for Complexa. None declared (M. L.).

Role of sponsors: The sponsor had no role in the design of the study, the collection and analysis of the data, or the preparation of the manuscript.

Other contributions: We thank Dinesh Khanna, MD, University of Michigan, for consultation on Delphi methodology and moderation of the face-to-face panel meeting, which was funded by Actelion Pharmaceuticals. We also thank Kelly Chin, MD, University of Texas, Southwestern Medical Center, for providing her expert opinion to survey questions and Stephen Chan, MD, PhD, University of Pittsburgh School of Medicine, for providing his expert opinion to survey questions and at the face-to-face PIXEL meeting. Medical writing support, funded by Actelion Pharmaceuticals, was provided by Holly Strausbaugh, PhD, and Laura Evans, PharmD, on behalf of Twist Medical.

Additional information: The e-Appendix and e-Figure can be found in the

Supplemental Materials section of the online article.

References

- Galiè N, Channick RN, Frantz RP, et al. Risk stratification and medical therapy of pulmonary arterial hypertension. *Eur Respir J*. 2019;53(1):1801889.
- Klinger JR, Elliott CG, Levine DJ, et al. Therapy for pulmonary arterial hypertension in adults: update of the CHEST guideline and expert panel report. *Chest*. 2019;155(3):565-586.
- Galiè N, Humbert M, Vachiery JL, et al. 2015 ESC/ERS guidelines for the diagnosis and treatment of pulmonary hypertension. *Eur Heart J*. 2016;37(1):67-119.
- Simonneau G, Torbicki A, Hoeper MM, et al. Selexipag: an oral, selective prostacyclin receptor agonist for the treatment of pulmonary arterial hypertension. *Eur Respir J*. 2012;40(4):874-880.
- Tapson VF, Torres F, Kermeen F, et al. Oral treprostinil for the treatment of pulmonary arterial hypertension in patients on background endothelin receptor antagonist and/or phosphodiesterase type 5 inhibitor therapy (the FREEDOM-C study): a randomized controlled trial. *Chest*. 2012;142(6):1383-1390.
- Jing ZC, Parikh K, Pulido T, et al. Efficacy and safety of oral treprostinil monotherapy for the treatment of pulmonary arterial hypertension: a randomized, controlled trial. *Circulation*. 2013;127(5):624-633.
- Tapson VF, Jing ZC, Xu KF, et al. Oral treprostinil for the treatment of pulmonary arterial hypertension in patients receiving background endothelin receptor antagonist and phosphodiesterase type 5 inhibitor therapy (the FREEDOM-C2 study): a randomized controlled trial. *Chest*. 2013;144(3):952-958.
- Sitbon O, Channick R, Chin KM, et al. Selexipag for the treatment of pulmonary arterial hypertension. *N Engl J Med*. 2015;373(26):2522-2533.
- Fitch K, Bernstein SJ, Burnand B, et al. *The RAND/UCLA Appropriateness Method User's Manual*. Santa Monica, CA: RAND; 2001.
- Ghio S, Klersy C, Magrini G, et al. Prognostic relevance of the echocardiographic assessment of right ventricular function in patients with idiopathic pulmonary arterial hypertension. *Int J Cardiol*. 2010;140(3):272-278.
- Raymond RJ, Hinderliter AL, Willis PW, et al. Echocardiographic predictors of adverse outcomes in primary pulmonary hypertension. *J Am Coll Cardiol*. 2002;39(7):1214-1219.

12. Sachdev A, Villarraga HR, Frantz RP, et al. Right ventricular strain for prediction of survival in patients with pulmonary arterial hypertension. *Chest*. 2011;139(6):1299-1309.
13. van Wolferen SA, Marcus JT, Boonstra A, et al. Prognostic value of right ventricular mass, volume, and function in idiopathic pulmonary arterial hypertension. *Eur Heart J*. 2007;28(10):1250-1257.
14. Frost AE, Badesch DB, Miller DP, et al. Evaluation of the predictive value of a clinical worsening definition using 2-year outcomes in patients with pulmonary arterial hypertension: a REVEAL Registry analysis. *Chest*. 2013;144(5):1521-1529.
15. Rhee RL, Gabler NB, Praetgaard A, Merkel PA, Kawut SM. Adverse events in connective tissue disease-associated pulmonary arterial hypertension. *Arthritis Rheumatol*. 2015;67(9):2457-2465.
16. Rubenfire M, Huffman MD, Krishnan S, et al. Survival in systemic sclerosis with pulmonary arterial hypertension has not improved in the modern era. *Chest*. 2013;144(4):1282-1290.
17. Chung L, Farber HW, Benza R, et al. Unique predictors of mortality in patients with pulmonary arterial hypertension associated with systemic sclerosis in the REVEAL registry. *Chest*. 2014;146(6):1494-1504.
18. National Institutes of Health Clinical Center. Phase III clinical worsening study of UT-15C in subjects with PAH receiving background oral monotherapy (FREEDOM-EV). NCT01560624. [ClinicalTrials.gov](https://clinicaltrials.gov/ct2/show/NCT01560624). Bethesda, MD: National Institutes of Health; 2012. <https://clinicaltrials.gov/ct2/show/NCT01560624>. Updated May 7, 2019.
19. Taichman DB, McGoon MD, Harhay MO, et al. Wide variation in clinicians' assessment of New York Heart Association/World Health Organization functional class in patients with pulmonary arterial hypertension. *Mayo Clin Proc*. 2009;84(7):586-592.