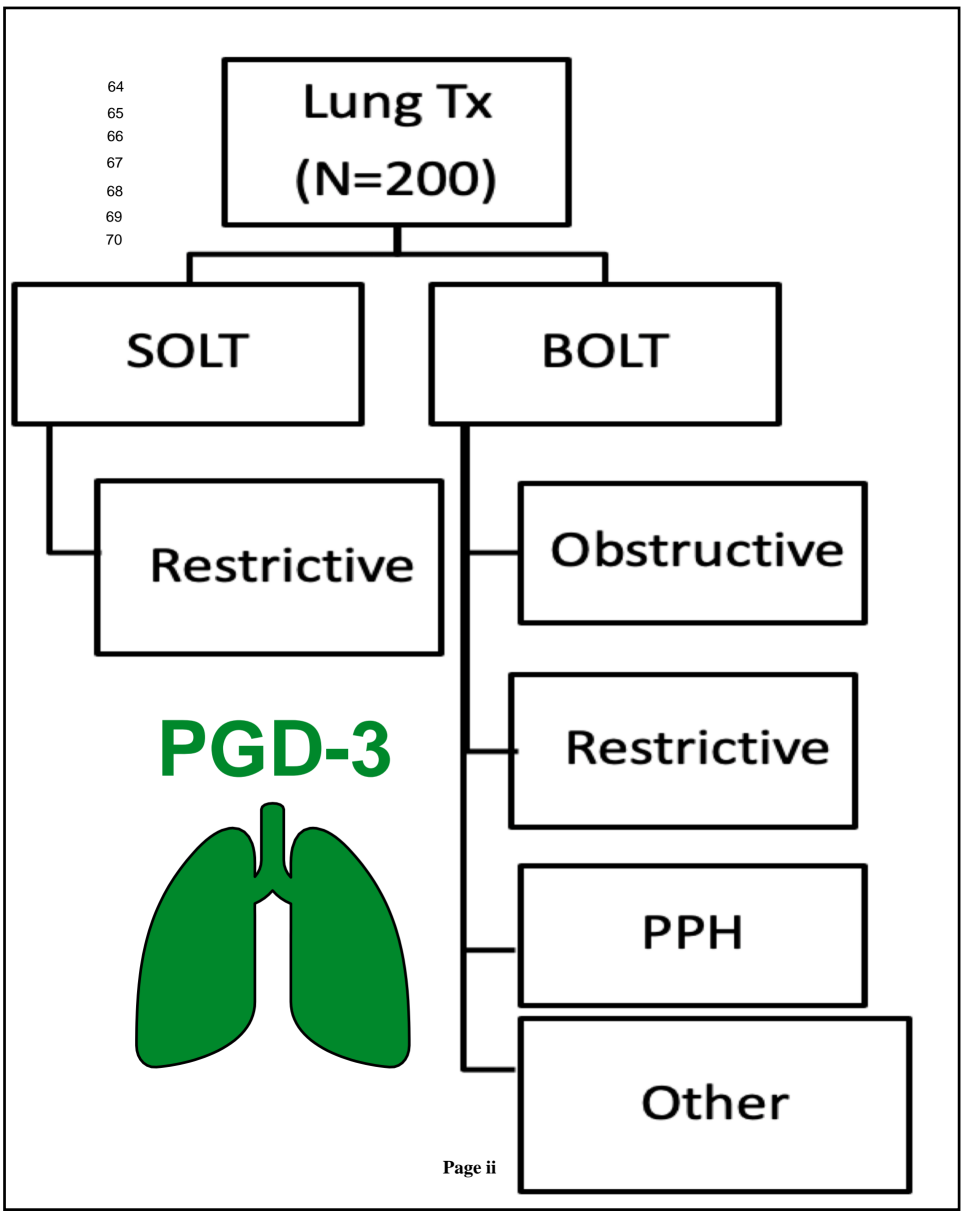
# Original iPVD Trial Protocol and Protocols for Lung Transplant Participants

Table of Contents	Page(s)
Randomization strata for Lung Transplants	ii
Original trial protocol (edited May 2017) - includes Lung Transplant and Advanced Heart Failure populations	1-6
Original clinicaltrials.gov page for NCT03081052 - includes Lung Transplant and Advanced Heart Failure populations	7-10
Protocol - Adult Mechanical Ventilation	11-19
Protocol – Epoprostenol: Adult Inhaled Pulmonary Vasodilator	20-31
Protocol - Transport of Mechanically Ventilated Cardiothoracic Patient on iNO or iEPO	
Protocol – OR support for Epoprostenol	
Protocol – Return to OR from Cardiothoracic ICU on Epoprostenol	37
Protocol – Inhaled Nitric Oxide for the Adult Lung Transplant patients in the Cardiothoracic ICU	38-41
Protocol – Inhaled Epoprostenol for the Adult Lung Transplant Patient in the Cardiothoracic ICU	42-45
Final trial protocol (edited March 2019) - includes Lung Transplant and Advanced Heart Failure populations	46-56



**1. Protocol Title:** Inhaled Pulmonary Vasodilator
Therapy in Left Ventricular Assist Device (LVAD)
Implantation, Heart Transplantation, and Lung
Transplantation: Prospective, Randomized,
Double-Blinded Study

## 77 2. Purpose of the Study:

71

78 1. Aim I - Clinical Trial Investigation. In order to 79 utilize Inhaled Epoprostrenol (iEPO, Veletri®, 80 Actelion Pharmaceuticals, South San Francisco, 81 CA, USA) as an acceptable alternative to Nitric 82 INOMAX®, Oxide (iNO, Mallinkrodt 83 Pharmaceuticals, St. Louis, MO, USA) in adult 84 patients, we propose a randomized, prospective, double-blinded trial in the cardiothoracic surgical 85 86 population, which will evaluate the primary 87 hypothesis that these two medications will have 88 similar efficacy in pulmonary vasodilation and a 89 similar impact on clinical outcomes in end-stage 90 lung disease patients undergoing lung transplantation and end-stage heart failure patients under durable LVAD implantation or 91 92 heart transplantation (Table 1). 93

94 2. Aim II – Cost-Capture Analysis. There will be a
95 parallel prospective cost-capture analysis
96 designed to precisely acquire the expenses that
97 each drug incurs per patient averaged across all
98 patients randomized to that drug.

§Sample Size	N = 424 (50/50 by randomization strata)
Population	1. Lung transplantation (N = 200)
	2. Heart Transplantation / LVAD implantation (N = 224)
Rationale	<ul> <li>Comparison of iNQ and iEpo impact on outcomes –</li> </ul>
	evaluate for equivalency
	<ul> <li>PVD therapy indications:</li> </ul>
	<ol> <li>Lung Transplant: Improvement of ventilaton and</li> </ol>
	perfusion matching after lung allograft
	implantation by vasodilation of ventilated
	pulmonary capillaries
	<ol><li>Heart Transplant/LVAD Implantation:</li></ol>
	Improvement of RV contractility after cardiac
	allograft implantation or LVAD implantation by
	PVR reduction
Study Design	Prospective, Randomized, Double-Blinded
Primary	<ol> <li>Lung Transplant: Severe PGD (grade 3)</li> </ol>
Outcomes	2a. Heart Transplant: RVAD insertion
	b. LVAD: INTERMACS Moderate or Severe RVF
Secondary	All populations:
Outcomes	<ul> <li>ICU LOS (days)</li> </ul>
	<ul> <li>Hospital LOS (days)</li> </ul>
	<ul> <li>Mechanical ventilator duration (hours)</li> </ul>
	Postoperative AKI
	<ul> <li>In-hospital mortality</li> </ul>
	<ul> <li>Mortality 30-day, 90-day, 1-year</li> </ul>
Study length*	24-36 months
	d for primary outcomes

Study length is determined through sample size divided by annual operations at Duke University Hospital; AKI = Acute kidney injury; iEpo = inhaled enoprostenol; iNO = Inhaled nitric oxide; ICU = Intensive care unit; LOS = Length of stay; LVAD = Left ventricular assist device; PGD = Primary graft dysfunction; PVD = Pulmonary vasodilators; PVR = Pulmonary vascular resistance; RVF = Right ventricular failure

## 99 3. Background & Significance:

00 Introduction. Inhaled Nitric Oxide (iNO) is a selective pulmonary vasodilator (PVD) with FDA-approval in the neonatal population alone. In adult patients, iNO is used off-label to treat pulmonary hypertension, right 01 ventricular (RV) failure, and ventilation-to-perfusion mismatch. Adult patients who undergo durable LVAD 02 03 implantation (e.g. Heartware®, Heartmate 2®, or Heartmate 3®) or cardiac transplantation for end-stage heart 04 failure or those that have endured lung transplantation as a result of end-stage lung disease, compose the largest subpopulation which receives PVD therapy at Duke University Hospital. Intravenous Epoprostenol is 05 FDA approved for adult patients with pulmonary hypertension and is the only agent which has displayed 06 mortality benefit in these patients. The inhaled formulation of Epoprostenol (iEPO) was developed in order to 07 08 maintain efficacy and avoid the systemic side effects of vasodilation and thrombocytopenia. Inhaled iEPO is 09 used off-label in our cardiothoracic surgical patients for new-onset perioperative pulmonary arterial 10 hypertension (PAH), known preoperative PAH, RV dysfunction with LVEF > 35-40%, and promotion of ventilation to perfusion matching through alveolar deposition of the prostanoid compound and vasodilation of 11 12 the intimately associated intra-acinar pulmonary arteries. This vasodilation can decrease pulmonary vascular 13 resistance and can improve oxygenation while avoiding systemic effects commonly seen in the intravenous 14 formulation. iEPO has been introduced in the cardiothoracic operating rooms (OR) and ICU as a cost-15 conscious alternative medication to iNO. iEPO may display an equivalent efficacy profile to iNO for pulmonary vasodilation and oxygenation and have a similar impact on clinical outcomes. For the purposes of this writing, 16 17 thoracic transplantation will refer to both heart and lung transplantation.

Pharmacology. There are 3 major pathways that affect pulmonary vascular tone: 1)Nitric oxide (vasodilatory), 18 2)Prostaglandin (vasodilatory), and 3)Endothelin (vasoconstrictive) pathways. During cardiothoracic 19 20 operations, particularly transplantation and LVAD surgery, there is an appreciable imbalance in these 21 pathways, which favors vasoconstriction. iNO administration, exerts its mechanism of pulmonary vasodilation 22 23 24 25 and ventilation-to-perfusion matching through exogenous NO delivery and iEPO applies a similar mechanism via exogenous prostacyclin delivery. Both agents are delivered through mechanical ventilation to ventilated alveoli in order to promote gas exchange at the capillary bed. Both inhaled medications are desirable in this population due to pulmonary selectivity, absence of systemic vasodilation, as well as fast onset (5-10 seconds 26 for iNO and 30-60 seconds for iEPO) and quick titration owing to short-half lives (10-20 seconds for iNO and 1-27 2 minutes for iEPO). There is no decision tree involved in the use of iNO vs iEPO except for that patient's 28 known drug allergies which may preclude use of one inhaled agent in favor of the other. Of note, endothelin 29 antagonists (e.g. bosentan), which are not part of our perioperative standard practice, are PO medications 30

#### Version 05/01/2017

33

34 which require reliable gastrointestinal absorption that may not be present during high-dose inotropic support, 35 and are not readily titrated to effect as are the inhaled PVD, iNO and iEPO.

36 Contraindications and Adverse Effects. Absolute indications for iNO in favor of iEPO are due to prostaglandin 37 allergy leading to anaphylaxis (extremely rare) or if the patient is pregnant due to risk for labor induction as a 38 result of prostacyclin agonism. Routine pregnancy testing is performed in the preoperative setting in line with 39 established preoperative anesthesia testing criteria. Parturients rarely present for thoracic transplantation or 40 LVAD implantation. There are no absolute contraindications to iNO therapy in adult patients but the iNO 41 delivery device system routinely measures the toxic metabolite of iNO, nitrogen dioxide (NO<sub>2</sub>), which can lead to hypoxemia during metabolite accumulation. Additionally, methemoglobinemia (MetHb) is another rare 42 43 adverse occurrence of prolonged iNO administration and MetHb levels are measured during arterial blood gas 44 analysis.

45 Preliminary retrospective study supporting noninferiority hypothesis. In a retrospective study of 51 adult 46 cardiothoracic surgical patients (all-comers, including thoracic transplantation, durable LVAD implantation, and 47 non-transplant and non-LVAD cardiac surgical patients), requiring pulmonary vasodilation, our group illustrated 48 similar efficacy between the use of iEPO and iNO with respect to optimizing RV hemodynamic variables,

49 including pulmonary vasodilation and mixed 50 venous oxygenation (Table 2). During this 51 investigation, iNO was initiated in the 52 operating room (OR) and continued during 53 transport and into the ICU. While in the ICU, 54 postoperative hemodynamic stability was 55 achieved within 2 hours and iNO was 56 transitioned to iEPO over 30 minutes in order

Table 2. Hemodynamic values in CT surgical patients comparing inhaled Nitric Oxide and Epoprostenol										
N= 51	*HR	<sup>¶</sup> MAP	<sup>¶</sup> PAPs	<sup>¶</sup> PAPd	<sup>¶</sup> PAPm	<b>ICVP</b>	*CI	PI	§LVAD flow	SVO2 (%)
<sup>a</sup> iNO	98	78	37.9	18.6	25.3	12.5	2.61	5.36	4.66	71
aiEpo	100	80	39.1	19.0	26.8	12.2	2.67	4.93	4.82	70
P-value	0.41	0.40	0.48	0.58	0.24	0.74	0.63	0.52	0.65	0.52

a = reported as mean values; \* units = beats per minute; ¶ units = mm Hg; ¥ units = L/min/m<sup>2</sup> CI = Cardiac Index, CVP = Central Venous Pressure, HR = Heart Rate, iNO = Inhaled nitric oxide, iEpo = Inhaled epoprostenol, LVAD = Left Ventricular Assist Device, MAP = Mean Arterial Pressure, PAPs = Systolic Pulmonary Artery Pressure, PAPm = Mean Pulmonary Artery Pressure, PAPd = Diastolic Pulmonary Artery Pressure, PI = Pulsatility Index, SvO<sub>2</sub> = Mixed Venous Oxygen Saturation

57 to provide continuous inhaled pulmonary vasodilation and allow the patient to self-control during medication 58 cross-over between iNO and iEPO. Clinical variables were followed at 5-minute intervals for 1 hour after 59 transition to iEPO. No statistically significant differences were seen in hemodynamic variables during this 60 transition (Table 2). The small sample size and retrospective design, however, incorporated several 61 confounding variables that could not be controlled and prospective data was deemed necessary to achieve 62 reliable conclusions by evaluating clinical outcomes in order to change clinician practice patterns. Other 63 investigations have demonstrated equivalence in hemodynamic variables, mixed venous oxygenation, and 64 ventilation-to-perfusion matching when delivery of iNO was compared with iEPO. These studies were, however, also retrospective or inadequately powered to rely on conclusions related to outcome measures. 65

66 The large cost differential between these two agents remains an important concern for the health system: iNO 67 is approximately 8-fold more expensive than iEPO, according to preliminary estimates based on PVD usage. Previous reports have estimated the cost of iNO administration to be between \$95.00 - \$115.00 per hour 68 69 during medication delivery. The cost, however, has not precisely captured the time required to assemble the iNO delivery system as well as resources utilized to breakdown this setup into individual components following 70 71 termination of delivery. The cost of iEPO delivery is captured at \$14.83 per hour, which includes solution 72 compounding by pharmacy as well as processing for delivery and nebulization by respiratory care services. 73 Additionally, the iEPO delivery-system setup is a one-time, fixed cost for the duration of administration. Similar 74 secondary resource utilization capture for iEPO is required for accurate cost comparison between these two 75 agents.

#### 76 4. Design & Procedures:

#### 77 <u>Aim I – Development of a Definitive Clinical Trial Investigation.</u>

78 1. Randomization and Double-Blinding. The clinical research unit (CRU) will receive preoperative notification of 79 lung and heart transplantation patients by reviewing the transplant waitlist. Preoperative notification of LVAD 80 implantation will be done by the review of the cardiothoracic surgical schedule. Using a 50% randomization process utilized and established by the CRU at Duke University Hospital, each eligible patient will be 81 82 randomized to receive either iNO or iEPO. The primary endpoint data will be collected and documented in an 83 electronic data capture system during the period of time the patient, clinical care team, and study team are 84 blinded. Primary endpoint data collection will be complete prior to the subjects' discharge from the ICU, at 85 which point the unblinding will occur. Since primary endpoint data collection will occur during the blinded period, the potential for bias will be substantially minimized. 86

2. Measured Outcomes. The primary endpoint for the comparison of efficacy in the Lung Transplant population 87 will be the incidence of Grade 3 Primary Graft Dysfunction (PGD). This is defined by the International Society 88 89 of Heart and Lung Transplantation (ISHLT) as severe hypoxemia with a PaO<sub>2</sub>-to-FiO<sub>2</sub> ratio < 200 or the 90 presence of venovenous extracorporeal membrane oxygenation (VV ECMO) at an time-point within the first 72

91

92 hours after lung transplantation. The primary endpoint for the comparison of efficacy in LVAD patients will be 93 incidence of moderate or severe RV failure according to Interagency Registry for Mechanically Assisted 94 Circulatory Support (INTERMACS) scoring. The primary endpoint for the comparison of efficacy in the heart 95 transplant subset will be the incidence rate of RVAD insertion. Secondary endpoints related to clinical 96 outcomes for all populations will be duration of postoperative mechanical ventilation, , ICU Length of Stay 97 (LOS), hospital LOS, incidence of acute kidney injury, incidence of in-hospital mortality, as well as 98 postoperative mortality at 30-days, 90-days, and 1-year after operation (Table 1).

#### 99 Aim II – Cost-Capture Analysis.

00 In parallel with the design & procedures of Aim I, the cost capture analysis component will be essential in order 01 to better gauge the cost due to duration of administration (variable cost) according to each inhaled PVD. 02 Established clinical criteria specific to each group (lung transplantation vs. heart transplantation/LVAD 03 implantation) have been developed to determine the inception of protocolized PVD weaning. Weaning 04 medications according to established protocols will allow for accurate interpretation of the comparative length 05 of therapy between iNO and iEPO and help prevent erroneous PVD usage without criteria for discontinuation. 06 Secondary resource utilization will be documented by respiratory care services and itemized cost sheets will be 07 developed.

#### 08 Subject Groups

09 Inhaled PVD therapy is administered to every patient undergoing thoracic transplantation and LVAD 10 implantation at our institution and each patient is eligible for enrollment. Over a 3-year period (1 year for followup) we will prospectively enroll 200 lung transplant subjects and 224 heart transplant or LVAD implantation 11 12 patients who will be informed and consented prior to their scheduled procedure. Potential subjects will be 13 under the care of 1 or more investigators in this study. Consented subjects will be randomly assigned to 1 of 2 14 groups, iNO vs iEPO, to be initiated in the OR on the day of the operation based on accepted standard of 15 practice and study protocol. Medication administration will be double-blinded, such that neither the surgical nor 16 anesthesiology teams will be notified of the inhaled agent to which the patient has been randomized. Ability to 17 unblind the delivery system will be made available to both teams if required to preserve optimal patient care. 18 As per our standard practice, respiratory care services will manage the initiation and maintenance of inhaled 19 PVDs in the OR and ICU, and these personnel will be the only practitioners notified of the actual delivered 20 medication during study blinding.

# 21 **Exclusion Criteria** 22 23 24 25 26 27 28

- Combined Organ Transplantation (Heart-Lung, Heart-Liver, Heart-Kidney)
- Age < 18 years old
  - Pregnancy (females of child bearing potential will receive pregnancy testing prior to cardiothoracic surgery as a standard of care)
  - Known allergy to prostaglandin (rare)
  - Subject is enrolled in another study protocol, which does not allow randomization of PVD therapy
  - Heart transplant or durable LVAD recipients with adult congenital heart disease (CHD)
  - Caveat: Does NOT meet exclusion criteria if the scheduled heart transplant or LVAD implantation is due to heart failure from a previous heart transplantation related to CHD, performed more than 90 days previous to the date of trial enrollment
  - Patient is scheduled to undergo lung transplantation but has undergone heart transplantation in the previous 90 days
  - Patient is scheduled to undergo durable LVAD implantation but has undergone heart transplantation in the previous 90 days
  - Patient is scheduled to undergo heart transplantation but has undergone lung transplantation in the previous 90 davs
  - Patients with preoperative VV ECMO as a bridge to lung transplantation

29

34

35

36

37

40	Stopping Criteria – In the event the following criteria are met and the clinical team is in agreement, subjects will
41	be weaned off of their iPVD per instutional standard iPVD weaning practice. If adverse events are
42	encountered, the drug will be immediately stopped without weaning.
43	Venoarterial (VA) ECMO insertion remains at end of operation
44	VA ECMO insertion is performed postoperatively in the ICU

- VA ECMO insertion is performed postoperatively in the ICU
- 45 LVEF < 30% on echocardiogram at the end of the operation for heart and lung transplant subjects
- 46 LVEF < 30% for heart and lung transplant subjects on echocardiogram noted postoperative in the ICU

47 48

49

50

- Inhaled pulmonary vasodilation is halted for reasons other than standard weaning ordered by the clinical care team
  - Adverse events related to the INO or EPO that affect the subject's welfare

### 51 Data Collection

52 Secondary measures will be hemodynamic variables (similar to those measured in Table 2) such as 53 transesophageal echocardiographic (TEE) evaluation of RV function based on stand-of-practice protocol, 54 intravenous administration of inotropes, serial measures of postoperative serum creatinine and GFR, resolution 55 of elevated liver function tests (heart failure patients, illustrates improvement in RV function), incidence of 56 thrombocytopenia (platelet count < 150 x  $10^{9}$ /L) and trajectory of resolution, as well as ventilation-to-perfusion 57 matching (arterial oxygen tension, PaO<sub>2</sub>; arterial carbon dioxide tension, PaCO<sub>2</sub>; and fraction of inspired 58 oxygen, FiO<sub>2</sub>). Variables will be recorded at designated time points during the entire duration of administration 59 – from initiation in the operating room to cessation in the ICU. These time points include: Intraoperative before 60 surgical incision, time = 0 (initiation of PVD), 30 minutes, 2 hours, 6 hours, 12 hours, 18 hours, 24 hours, and 61 every 6 hours up through 72 hours after initiation. These secondary measures will be obtained up through 72 hours after initiation regardless of cessation or continuation of the inhaled PVD. After 72 hours, increments of 62 63 every 12 hours thereafter will be assessed if PVD administration continues. Ventilation and perfusion nuclear 64 scans will be obtained and recorded per standard clinical practice for each group of lung transplant recipients. 65 Established protocols with criteria for initiation of medication weaning have been created according to each 66 medication based on individual pharmacokinetic properties. Once established criteria are met, weaning of each 67 inhaled PVD will begin and continue until the medication is terminated according to standardized weaning 68 protocols established for lung transplant patients and heart transplant/LVAD patients. 69

Subject follow up. Subject will be contacted by phone by a member of the research team and be asked a short series of questions to assess their current medical condition and any changes since surgery at 30-days (± 3 days), 90-days (± 5 days), and 1-year (± 7 days) after surgery completion date. The phone follow-up should take approximately 5 minutes of the subject's time. If subjects have been admitted to a hospital outside of Duke Health after surgery they will be asked to sign an authorization of release to provide us permission to obtain medical information related to their hospitalization.

## 76 Blood Sampling

77 Blood samples will be drawn for analysis as a part of this study. One 9 ml sample of blood will be obtained 78 from each patient prior to the initiation of PVD therapy and stored at 4°C prior to processing. This sample will 79 be stored for Genomic DNA analysis at the completion of this study in order to assess patients who are 80 responders to inhaled pulmonary vasodilaton through upregulation and down regulation of notable vasoactive 81 substances (e.g. endothelin, thromboxane, nitric oxide, prostaglandin, etc.). In addition, each subject will also 82 be asked to sign the Genomic and Proteomic Database Repository (IRB Pro00015651) consent form, thus 83 84 85 86 87 allowing the banking of their plasma and DNA samples as well as data to be used for future research. Participation in IRB Pro00015651 is voluntary and optional to all subjects consented in this parent study. Blood samples (7 ml each) will be drawn at 3 separate time points: 1) directly after insertion of the invasive blood pressure monitoring (arterial) line, 2) POD 1, and 3) POD 7. In each 7ml blood sample, 3.5ml will be collected in Sodium Citrate tubes for coagulation analysis and another 3.5ml will be collected in EDTA tubes 88 for metabolomic and proteomic analysis. Plasma will be separated from these samples and banked at -80°C for analyses of proteomic and metabolomic signatures. Up to 30ml of blood will be collected during the 12 89 90 month study participation period.

91

92 6. Subject Identification, Recruitment, & Compensation: Subjects will be recruited either during the 93 outpatient or inpatient evaluation phase, or contacted by phone. Recruitment may also occur on the day of the operation given the complexities of the transplant process, which may provide obstacles to earlier enrollment. 94 95 After obtaining permission from the operating surgeon, surgical subjects will be screened by the study 96 coordinator by reviewing the transplant pre-list. Prior to asking any patient for consent to participate, the patient 97 or Legally Authorized Representative (LAR) will be approached first by the surgeon or one of the members of 98 the surgical care team to determine if the patient or LAR is willing to consider enrollment in the study. If so, the 99 subject or LAR will either be seen during an inpatient or outpatient visit, or be contacted by phone and 00 informed about the study by a member of the research team. If the individual or LAR is willing to consider 01 enrollment and does not meet exclusion criteria, then the research coordinator will present the research 02 protocol in its entirety. During this time, the study coordinator will answer any and all questions as they arise. If 03 the subject or LAR agrees to participate, the coordinator will ask the them to sign and date the appropriate 04 consent form. A copy of this consent form will be given to the subject and a copy of the consent form will be

- added to the subject's medical record. The subject or LAR will be given the option to sign a separate consent form to allow us to store portions of the collected blood specimens and any data collected under this research study and maintain these samples and data in a database/repository (PRO00015651) for possible use in future research studies relating to surgical outcomes. In the event a LAR provides consent at the time of enrollment, the subject will be approached once they regain the ability to provide an informed consent.
- 11 Recruitment will not routinely occur on the day of the operation and most patients will be enrolled at least 12 12 hours in advance and provided at least the allowable time to review the study consent form and discuss their 13 options with the PI and study personnel. There will be no direct compensation to the patient for recruitment.
- If a subject is enrolled and randomized in this study for their LVAD implantation procedure and is later planned
   to receive a heart transplant, that previously enrolled subject is eligible to be re-enrolled. The following caveats
   apply to this subpopulation of LVAD patients:
- 17 A)Durable LVAD implantation may occur as a bridge to heart transplantation.
- B)If LVAD implantation is followed by heart transplantation WITHIN 1 year following LVAD implantation, then
   data collected up through the time of heart transplantation will be recorded and valid as a patient in the LVAD
   group.
- C)Data collected on or after the date of LVAD explantation/heart transplantation for such a patient will be considered as part of the heart transplant group.
- D)If LVAD implantation is followed by heart transplantation AFTER 1 year following LVAD implantation, then
   the 1year follow-up period is complete and the patient may re-enter the trial as a heart transplant patient.
- If a subject is enrolled and randomized in this study for their durable LVAD implantation procedure and is scheduled to receive a new durable LVAD via an LVAD exchange operation, the subject is eligible to be reenrolled.
- 28
   29 7. Subject's Capacity to Give Legally Effective Consent: Explicit (written) consent will be obtained from the patient or the patient's legal decision maker.
   31
- 8. Study Interventions: Using a 50% randomization process utilized and established by the CRU, each eligible patient will be randomized to receive either iNO or iEPO, to be initiated in the OR based on accepted standard of practice at Duke University Hospital, during the clinical care of these patients.
- 36 9. Risk/Benefit Assessment: There is no direct benefit of this study to the enrolled subjects. Data gathered 37 from this study may benefit future patients. Up to 30 ml of blood will be drawn during the 12 month study 38 participation period. Blood sampling will be obtained, in the majority of subjects, from indwelling arterial or 39 central venous lines inserted at the beginning of the intraoperative period as part of standard practice for these 40 operations and there will be no additional risk to the patient for obtaining such vascular access. On rare 41 occasion, blood sampling may be obtained from additional venipuncture sites during the postoperative period. 42 Risks of blood sampling if obtained through venipuncture are pain, swelling, possible infection at the site of 43 venipuncture. While these risks are minimal, the additional blood volume is highly unlikely to contribute to the patient's need for blood transfusion. To minimize any potential risk to the patient from genetic data, 44 investigators and patients will be blinded to the individual patient's genotype. This information will not be 45 46 included in the patient chart, will remain absolutely confidential, and will not be given to the patient or their 47 family. DNA samples will be identified only by a coded number whose relation to the patient's name and other identifiers is available only to the data manager. The identity of the patient will remain anonymous in any 48 49 publications which may result from this investigation. 50
- There will be no additional risks to the subjects as a result of this study. Prior to June of 2015, iNO was the sole option for inhaled pulmonary vasodilation in this patient population and therefore utilized in each operation for this indication. As of June 2015, iEPO was introduced for the same indications as iNO in order to serve as a cost-conscious alternative to iNO and to potentially explore a different, equally impactful pathway for clinically evident pulmonary vasodilation (as measured by Swan-Ganz catheter data and determined by transesophageal echocardiography). There are no additional risks to the patient aside from the rare adverse effects such as allergic reaction, as previously discussed. The most common side effect of iNO is hypotension. The side effects common to intravenous iEPO are nausea, vomiting, hypotension, flushing, chest pain, anxiety, dizziness, bradycardia, difficulty breathing, abdominal pain, musculoskeletal pain and tachycardia.
- 60 **10. Costs to the Subject:** There will be no additional costs to the subjects as a result of this study.

62 11. Data Analysis & Statistical Considerations: Summary statistics will be computed for demographic, 63 clinical, and outcome variables in the form of frequencies (percentage) for categorical variables and mean 64 (standard deviation) for continuous variables for each arm. Univariate analysis will be performed to compare 65 the difference of each variable between treatment groups by chi-square or Fisher exact tests for categorical 66 variables, and t-tests or Wilcoxon Rank-Sum tests for continuous variables depending on data normality. The 67 univariate results for the outcome variables will provide information on iNO treatment effect in comparison to 68 iEPO without taking into account other potential confounding factors. All non-outcome variables meeting p< 69 0.15 association with treatment group will be considered for variable selection to build a multivariable 70 regression model. For each outcome of interest, we will start with a regression model (logistic regression for 71 72 73 74 binary outcomes or generalized linear model for continuous outcomes) with all variables selected from univariate analysis described above. Based on stepwise variable selection, we will determine the final set of covariates to be included in the final multivariable model to test the treatment group effect. Based on the analysis results, we will be able to understand if iNO is equivalent to iEPO (no significant difference) or 75 significantly better or worse than iEPO (significant treatment effect) to address the efficacy of iNO for Aim 1. 76 Several of secondary measures will be obtained over time. We will apply generalized mixed model to take into 77 account the repeated measures over time to test for treatment effect. In the case of patients have switched to 78 the other arm due to clinical decision, we will conduct the primary analysis based on the intent to treat (ITT) 79 without reclassifying treatment assignment. In addition, protocol analysis, where only patients follow the 80 protocol assignment are included will also be conducted to verify ITT results. For Aim 2 to compare cost 81 capture analysis, the comparison of cost measures between two groups will be tested by two sample t-test.

82 Based on recent annual operations, approximately 120 LVAD implantations, 60 heart transplantations, and 110 83 Lung transplantations were performed at Duke University Hospital during FY 2014 - 2015. This study has been 84 individually powered to primary endpoints for each arm (Table 1) and the duration of study enrollment has 85 been determined according to annual operations and sample-size calculations. We estimated sample size 86 based on equivalence test of the incidence rates of a binary outcome (e.g. PGD grade 3 (PGD-3)) of two 87 treatment groups as an illustration. Assuming the incidence rate of PGD-3 under iEPO treatment is 0.35 and 88 acceptable margin of the equivalence is  $\pm$  0.19, we will need 224 patients to have 80% power to detect an 89 actual difference at 0.05 between two treatment group under this margin. This implies that the acceptable 90 range of incidence rate for iNO treatment is from 0.21 to 0.59. Based on this estimate, we propose to enroll 91 200 lung transplant patients and 224 LVAD and heart transplant patients (n = 424) over a period of 24 to 36 92 months; the exact time point for trial culmination between 24 and 36 months will be dependent on enrollment 93 rate. There will be a 50% randomization rate for each inhaled agent such that 212 patients will receive iEPO 94 and 212 patients will receive iNO. 95

96 12. Data & Safety Monitoring: The proposal is not introducing a new medication that has not been utilized by 97 our group and safety has been established for this patient population through clinical practice and medication 98 usage. Safety will, however, be determined by assessing reported, rare, adverse effects of iNO (systemic 99 hypotension, methemoglobinemia, and rebound pulmonary hypertension after appropriate weaning) and iEPO 00 (systemic hypotension, non-surgical bleeding related to thrombocytopenia, flushing, and rebound pulmonary 01 hypertension after appropriate weaning) in order to accurately monitor adverse events (AE) during this study. 02 The PI will review and sign off on AE's as they occur and perform a guarterly review and determine if AE's are 03 related to the study or otherwise. AE's will be reported to the IRB per HRPP policies.

04 *Stopping Rule:* Subjects who meet the stopping criteria in section 4 will continue to be enrolled and followed for 05 primary outcome analysis.

06 13. Privacy, Data Storage & Confidentiality: All data collected in the case report forms (CRF) will be collected by review of the subjects routine medical record documentation or during the intraoperative portion of 07 08 the study. All subjects will be given a study ID in an order to maintain their identity and subject's identity will be 09 protected and confidentially maintained. Barcodes will be affixed to each study sample collected according to 10 the protocol. For future review, the study number and barcode will be the only identifying information 11 associated with the subject. All paper data will be stored in a locked cabinet in the research teams office as 12 outlined in the research data security plan. Any computerized data will be stored within the Duke University 13 Medical Center's Database, which is password protected, and located behind Duke Computing firewalls. Only the PI and the statisticians will have access to the data obtained from these cases. 14



#### Try our beta test site

**IMPORTANT**: Listing of a study on this site does not reflect endorsement by the National Institutes of Health. Talk with a trusted healthcare professional before volunteering for a study. Read more...

Trial record 1 of 4 for: ghadimi
Previous Study | Return to List | Next Study

Inhaled Pulmonary Vasodilator Therapy in Left Ventricular Assist Device (LVAD) Implantation, Heart Transplantation, and Lung Transplantation: Prospective, Randomized, Double-Blinded Study (iNO vs iEPO)

This study is not yet open for participant recruitment. (see Contacts and L Verified March 2017 by Duke University Sponsor: Duke University Information provided by (Responsible Party): Duke University	439 440 441 442 443	ClinicalTrials.gov Identifier: NCT03081052 First received: March 8, 2017 Last updated: March 14, 2017 Last verified: March 2017 History of Changes
Full Text View         Tabular View         No Study Results Posted         Disclaiment	r How to	Read a Study Record

#### Purpose

The primary purposes of this study has 2 aims. First, to conduct a clinical investigation to determine if iEPO, Veletri® will have similar efficacy in pulmonary vasodilation and have a similar impact when compared to iNO, INOMAX® in adult patient who undergo a heart transplantation, lung transplantation or implantation of a left ventricular assist device. Second, to conduct a cost-capture analysis on the expense each drug incurs per patient.

Condition	Intervention	Phase
Heart Transplant Surgery	Drug: iNO	Phase 4
Lung Transplant Surgery	Drug: iEPO	

Study Type:	Interventional
Study Design:	Allocation: Randomized
	Intervention Model: Parallel Assignment
	Masking: Participant, Care Provider, Investigator
	Primary Purpose: Other

Official Title: Inhaled Pulmonary Vasodilator Therapy in Left Ventricular Assist Device (LVAD) Implantation, Heart Transplantation, and Lung Transplantation: Prospective, Randomized, Double-Blinded Study

#### Resource links provided by NLM:

MedlinePlus related topics: Heart Transplantation Lung Transplantation

Drug Information available for: Nitric oxide

U.S. FDA Resources

Further study details as provided by Duke University:

#### Primary Outcome Measures:

Incidence of Grade 3 Primary Graft Dysfunction (PGD) for Lung Transplant subjects. [Time Frame: Up to 72 hours ]

https://clinicaltrials.gov/ct2/show/NCT03081052?term=ghadimi&rank=1

Inhaled Pulmonary Vasodilator Therapy in Left Ventricular Assist Devic...andomized, Double-Blinded Study - Full Text View - Clinical Trials.gov

This is defined by the International Society of Heart and Lung Transplantation (ISHLT) as severe hypoxemia with a PaO2-to-FiO2 ratio < 200 or the presence of venovenous extracorporeal membrane oxygenation (VV ECMO) at an time-point within the first 72 hours after lung transplantation.

- Incidence of moderate or severe RV failure for the LVAD implantation subjects. [Time Frame: up to approximately 21 days after LVAD placement]
   This is defined by the Interagency Registry for Mechanically Assisted Circulatory Support (INTERMACS) scoring.
- Incidence of severe RV failure for Heart Transplantation subjects. [Time Frame: up to approximately 30 days after heart transplantation ]
   This is defined by the incidence of an RVAD placement

Secondary Outcome Measures:

- Duration of postoperative mechanical ventilation [Time Frame: up to approximately 90 days]
   Length of time from intubation until patient is extubated
- Per patient cost [ Time Frame: up to approximately 30 days ]
   Cost associated with the duration of PVD administration
- Length of ICU stay [ Time Frame: up to approximately 90 days ]
   Length of time from ICU admission from surgery until ICU discharge
- Length of hospital stay [ Time Frame: up to approximately 1 year ]
   Length of time from surgery to hospital discharge
- Incidence of Acute Kidney Injury [ Time Frame: up to approximately 14 days ] defined by KDIGO-AKI criteria
- Incidence of in-hospital mortality [Time Frame: up to approximately 1 year]
   Death that occurs during the hospital stay
- Incidence of postoperative mortality within 30 days [Time Frame: up to approximately 30 days ]
   From the day of surgery to 30 days (+/- 3 days)
- Incidence of post-operative mortality within 90 days [Time Frame: up to approximately 90 days ]
   From the day of surgery to 90 days (+/- 5 days)
- Incidence of post-operative mortality within 1 year [Time Frame: up to approximately 1 year ]
   From the day of surgery to 1 year (+/- 7 days)

Estimated Enrollment:424Anticipated Study Start Date:April 2017Estimated Study Completion Date:April 2021Estimated Primary Completion Date:April 2021 (Final data collection date for primary outcome measure)

Arms	Assigned Interventions
Active Comparator: Lung transplant with iNO	Drug: iNO Subject will receive inhaled Nitric Oxide in this intervention Other Name: Inhaled Nitric Oxide
Active Comparator: Lung transplant with iEPO	Drug: iEPO Subject will receive inhaled Epoprostrenol in this intervention Other Name: Inhaled Epoprostrenol
Active Comparator: Heart transplant & LVAD implantation with iNO	Drug: iNO

	Subject will receive inhaled Nitric Oxide in this intervention Other Name: Inhaled Nitric Oxide
Active Comparator: Heart transplant & LVAD implantation with iEPO	Drug: iEPO
	Subject will receive inhaled Epoprostrenol in this intervention
	Other Name: Inhaled Epoprostrenol

#### **Detailed Description:**

Inhaled Nitric Oxide (iNO) is a selective pulmonary vasodilator (PVD) with FDA-approval in the neonatal population alone. In adult patients, iNO is used offlabel to treat pulmonary hypertension, right ventricular (RV) failure, and ventilation-to-perfusion mismatch. Adult patients who undergo durable LVAD implantation (e.g. Heartware®, Heartmate 2®, or Heartmate 3®) or cardiac transplantation for end-stage heart failure or those that have endured lung transplantation as a result of end-stage lung disease, compose the largest subpopulation which receives PVD therapy at Duke University Hospital. Inhaled Epoprostenol (iEPO) has been introduced in the cardiothoracic operating rooms (OR) and ICU as a cost-conscious alternative medication to iNO. iEPO may display an equivalent efficacy profile to iNO for pulmonary vasodilation and oxygenation and have a similar impact on clinical outcomes.

424 informed and consented subjects undergoing thoracic transplantation or left ventricular assist device (LVAD) implantation under the care of one or more investigators will be prospectively enrolled over a three-year period (one-year for follow-up). Patients will be randomly assigned 50/50 according to randomization strata to one of two standard of care pulmonary vasodilation therapy, iNO vs iEPO. Additional study procedures will involve data collection and blood sampling.

#### Eligibility

Ages Eligible for Study:18 Years and older (Adult, Senior)Sexes Eligible for Study:AllAccepts Healthy Volunteers:No

#### Criteria

Inclusion Criteria:

- Thoracic (heart or lung) transplantation patients
- LVAD implantation patients

Exclusion Criteria:

- Combined Organ Transplantation (Heart-Lung, Heart-Liver, Heart-Kidney)
- Age < 18 years old
- Pregnancy
- Known allergy to prostaglandin (rare)
- Subject is enrolled in another study protocol, which does not allow randomization of PVD therapy
- Heart transplant or durable LVAD recipients with adult congenital heart disease (CHD) o Caveat: Does NOT meet exclusion criteria if the scheduled heart transplant or LVAD implantation is due to heart failure from a previous heart transplantation related to CHD, performed more than 90 days previous to the date of trialenrollment
- Patient is scheduled to undergo lung transplantation but has undergone heart transplantation in the previous 90 days
- Patient is scheduled to undergo durable LVAD implantation but has undergone heart transplantation in the previous 90 days
- Patient is scheduled to undergo heart transplantation but has undergone lung transplantation in the previous 90 days
- Patients with preoperative Venovenous ECMO as a bridge to lung transplantation

#### Contacts and Locations

Choosing to participate in a study is an important personal decision. Talk with your doctor and family members or friends about deciding to join a study. To learn more about this study, you or your doctor may contact the study research staff using the Contacts provided below. For general information, see Learn About Clinical Studies.

Please refer to this study by its ClinicalTrials.gov identifier: NCT03081052

#### Contacts

Contact: Tiffany L Bisanar, RN, BSN 919-681-0866 tiffany.bisanar@duke.edu Contact: Kamrouz **Ghadimi**, MD 919-681-6532 kamrouz.**ghadimi**@duke.edu Inhaled Pulmonary Vasodilator Therapy in Left Ventricular Assist Devic...andomized, Double-Blinded Study - Full Text View - Clinical Trials.gov

#### Sponsors and Collaborators

Duke University

#### Investigators

Principal Investigator: Kamrouz Ghadimi. MD Duke Health

## More Information

Responsible Party:	Duke University	,
ClinicalTrials.govIdentifier:	NCT03081052	History of Changes
Other Study ID Numbers:	Pro00078035	
Study First Received:	March 8, 2017	
Last Updated:	March 14, 2017	
Individual Participant Data		
Plan to Share IPD:	No	

Studies a U.S. FDA-regulated Drug Product:	Yes
Studies a U.S. FDA-regulated Device Product:	No
Product Manufactured in and Exported from the U.S.:	No

Keywords provided by Duke University: Heart and Lung transplantation surgery Pulmonary vasodilation therapy

Additional relevant MeSH terms: Nitric Oxide Vasodilator Agents Bronchodilator Agents Autonomic Agents Peripheral Nervous System Agents Physiological Effects of Drugs Anti-Asthmatic Agents Respiratory System Agents

ClinicalTrials.gov processed this record on May 05, 2017

Free Radical Scavengers Antioxidants Molecular Mechanisms of Pharmacological Action Neurotransmitter Agents Endothelium-Dependent Relaxing Factors Gasotransmitters Protective Agents

574					
575 576 577	Adult Mechanical Ventilation Protocol for Routine Post-Op Cardiothoracic Surgery and Lung Transplant Patients				
578					
579	Purpose: To provide consistent clinical practice and timely interventions in the management of				
580	cardiothoracic surgical (CTICU) patients who require mechanical ventilation.				
581					
582					
583	Protocol Initiation:				
584					
585	The Mechanical Ventilation Protocol will be initiated upon receipt of a provider's order e.g. "CTICU				
586	Mechanical Ventilation Protocol".				
587					
588	Upon receipt of a provider's order, the respiratory care practitioner will:				
589					
590	Based upon the patient's diagnosis and surgical procedure, select the appropriate arm of the				
591	protocol from the following choices:				
592	<ul> <li>Ventilation Protocol - Cardiothoracic Surgery Patients (Section IV)</li> </ul>				
593	<ul> <li>Ventilation Protocol - Lung Transplant Surgery Patients (Section V)</li> </ul>				
594	Determine initial ventilator settings and initiate mechanical ventilation.				

595 596	<ul> <li>Assess the patient's response to the level of mechanical ventilation support provided.</li> <li>Communicate the initial mechanical ventilation settings and all subsequent changes to other</li> </ul>				
597 598	<ul> <li>members of the patient care team.</li> <li>Document all interventions in the medical record.</li> </ul>				
599					
600	Ventilation Protocol – Cardiothoracic Surgery Patients				
601	Goals				
602	$\Box$ To maintain the patient's arterial pH between 7.35 and 7.45				
603	To maintain the patient's PaO2 between 60 and 85 mm Hg				
604	· •	To maintain the patient's PaCO2 < 50 mm Hg			
605	$\Box$ To maintain the patient's SpO2>90%				
606	□ Tomaintain an ETCO2 < 45 mm Hg and PaCO2 < 50 mm Hg (during re-warming and shivering)				
607	$\Box$ To provide an inspiratory pressure plateau of no greater than 30 cm H2O				
608					
609	Initial Mode and Setting Selection				
610					
611	Mode: The initial mode may be one of the following:				
612	<ul> <li>Pressure Assist-Control (PAC)</li> </ul>				
613	<ul> <li>Pressure Support (PSV)</li> </ul>				
614	Inspiratory Pressure (Tidal Volume) - Adjusted to achieve targeted tidal volume and comfort				
615	<ul> <li>Inspiratory pressure will be set to achieve an exhaled VT of 4-8 mL/kg of the patient's</li> </ul>				
616	ideal bodyweight				
617	• Total pressure will not exceed 30 cm H2O.				
618	$\square Respiratory Rate (f) - 14 (then adjusted to control PaCO2)$				
619	□ Inspiratory Time will be set to optimize patient comfort, avoid or minimize air trapping, and				
620	produce an I:E ratio of less than 1:1				
621	□ FiO2 and PEEP				
622 623	<ul> <li>FiO2-0.6 (unless otherwise ordered by an esthesiologist)</li> <li>PEEP-5 cm H2O</li> </ul>				
624	<ul> <li>Pulmonary Vasodilator Therapy: Perprovider order.</li> </ul>				
625	<ul> <li>Non-invasive Monitoring (Continuous):</li> </ul>				
626	<ul> <li>Pulse Oximetry</li> </ul>				
627	<ul> <li>Capnography</li> </ul>				
628	<ul> <li>Continuously from admission to unit until extubation or for up to 24 hours po</li> </ul>	st			
629	admission.				
630	<ul> <li>Patients receiving trach collar trials.</li> </ul>				
631	<ul> <li>24 hours post reintubation</li> </ul>				
632	On the order of a provider				
633	Subsequent Adjustments/Weaning				
634	Obtain an ABG within 30 minutes after admission:				

page 12

635 636		<ul> <li>Assess the PaCO2-ETCO2 gradient. If &lt; 10 mm Hg, patient stable, no acidosis, use ETCO2 to adjust respiratory rate unless otherwise stated.</li> </ul>
637		Management during re-warming and shivering
638		<ul> <li>Adjust respiratory rate to keep ETCO2 &lt; 45 mm Hg, PaCO2 &lt; 50 mm Hg</li> </ul>
639		Oxygenation adjustments
640		• Wean FiO2
641		If SpO2 > 90% (PaO2 > 60 mm Hg) to a FiO2 goal of 0.4
642		<ul> <li>Notify MD if SpO2 &lt; 90%, PaO2 &lt; 60 mm Hg or if unable to obtain a FiO2 of 0.4</li> </ul>
643		within 4 hours.
644		<ul> <li>Increase FiO2/PEEP if indicated (see Appendix A)</li> </ul>
645		<ul> <li>For a patient with a VAD: Keep PEEP at 5 cm H2O unless directed otherwise by</li> </ul>
646		provider order.
647		Respiratory rate adjustments
648		• Patient must be hemodynamically stable without shivering or bleeding requiring
649 650		treatment, temperature > 36°C, responsive and breathing spontaneously (RN will begin
651		<ul> <li>weaning sedation when patient begins to waken.)</li> <li>Assess ABG to ensure PaO2 &gt; 60 mm Hg, PaCO2 &lt; 50 mm Hg, pH 7.35-7.45</li> </ul>
652		
653		Elevate HOB 30° unless otherwise ordered or contraindicated (i.e. intra-aortic balloon pump)
653 654		Change to PSV 10 cm H2O (40%, 5 PEEP) from PAC when ETCO2 < 45 mm Hg and reliable, spontaneous respiratory drive present.
655 656		<ul> <li>Extubation – Refer to the following</li> <li>Appendix D: Daily Spontaneous Breathing Assessment and Trial (SBT) and</li> </ul>
657		<ul> <li>Appendix D: Daily Spontaneous Dreating Assessment and that (SBT) and</li> <li>Appendix F: Extubation Criteria</li> </ul>
658		Weaningfrom Mechanical Ventilation – Trach Collar Trials
659		<ul> <li>Appendix E. Trach Collar Trial Weaning</li> </ul>
660		
661		
662	Ventila	tion Protocol - Lung Transplant Surgery Patients
663 664 665	Goals:	The goals of the Lung Transplant Surgery Protocol include all of the following:
666		To maintain the patient's arterial pH between 7.35 and 7.45
667		To maintain the patient's PaO2 between 60 and 85 mm Hg
668		To maintain the patient's SpO2>90%
669		To maintain the patient's PaCO2 < 50 mm Hg
670		To maintain an ETCO2 < 45 mm Hg and PaCO2 < 50 mm Hg (during re-warming and shivering)
671		Respiratory rate ≤ 16
672		To provide an inspiratory pressure plateau of no greater than 30 cm H2O
673	Initial	Ande and Setting Selection upon Admission to CTICU
674		Mode: The initial mode will be Pressure Assist-Control (PAC)
675		Inspiratory Pressure (Tidal Volume)
676		• Inspiratory pressure will be set to achieve an exhaled VT of 4-8 mL/kg of the patient's
677		ideal body weight.
678		<ul> <li>Total pressure will not exceed 30 cm H2O.</li> </ul>

679		Respiratory Rate (f) = 10 (then adjusted to control PaCO2)				
680		FiO2 and PEEP				
681		$\circ$ Start with a FiO2 of 0.21 to achieve a PaO2 > 65				
682		o PEEP=8cmH2O				
683		Inspiratory Time (Ti) will be set between 1.4-1.6 seconds on sedated patients.				
684		Pulmonary Vasodilator Therapy: Perprovider order.				
685		Non-invasive Monitoring (Continuous):				
686		<ul> <li>Pulse Oximetry</li> </ul>				
687		<ul> <li>Capnography</li> </ul>				
688		<ul> <li>Continuously from admission to unit until extubation or for up to 24 hours post</li> </ul>				
689		admission.				
690		<ul> <li>Patients receiving trach collar trials.</li> </ul>				
691		<ul> <li>24 hours post reintubation</li> </ul>				
692		<ul> <li>On the order of a provider.</li> </ul>				
693						
694	Subse	quent Adjustments and Weaning				
695						
696		Obtain an arterial blood gas within 30 minutes after admission:				
697		<ul> <li>Assess the PaCO2-ETCO2 gradient. If &lt; 10 mm Hg, patient stable, no acidosis, use</li> </ul>				
698		ETCO2 to adjust respiratory rate unless otherwise stated.				
699						
700		Oxygenation: : Appendix A: FiO2/PEEP Table				
701						
702		Management during re-warming and shivering				
703		$\circ$ Adjust respiratory rate to keep ETCO2 < 45 mm Hg, PaCO2 < 50 mm Hg				
704		<ul> <li>Treat shivering (nursing)</li> </ul>				
705		Elevate HOB 30° unless otherwise ordered				
706						
707		Weaning				
708		<ul> <li>Patient must be hemodynamically stable.</li> </ul>				
709		<ul> <li>Assess ABG to ensure PaO2 &gt; 65 mm Hg, PaCO2 &lt; 50 mm Hg, pH 7.35-7.45</li> </ul>				
710		$\circ$ Change to PSV 10 cm H2O (40%, 5 PEEP) from PAC when ETCO2 < 45 mm Hg, and				
711		reliable, spontaneous respiratory drive present.				
712	<b>–</b>					
713	Extuba	ation – Refer to the following				
714	_					
715		Appendix D: Daily Spontaneous Breathing Assessment and Trial (SBT) and				
716		Appendix F: Extubation Criteria				
717						
718	weani	ng from Mechanical Ventilation – Trach Collar Trials				
719	_					
720		Appendix E. Trach Trial Weaning				
721						
722						
723	Patien	t Assessment and Ventilator Monitoring – Patient assessment and ventilator monitoring will be				
724		ned to determine the patient's clinical status and progress toward goals.				
· — ·	P 511011					

725		he RCP will assess the patient and monitor the ventilator
726		<ul> <li>Immediately after initiating mechanical ventilation</li> </ul>
727		<ul> <li>At 6-hour intervals (approximately) thereafter</li> </ul>
728		<ul> <li>Whenever there is a change in the level of support (mode) provided or a change in</li> </ul>
729		settings that effects minute ventilation or mean airway pressure.
730		<ul> <li>Whenever there is an acute change in the patient's condition signaled by a rapid</li> </ul>
731		deterioration in vital signs or oxygenation or a change in ventilation.
732		atient assessment and ventilator monitoring will consist of
733		<ul> <li>An evaluation of the performance of the mechanical ventilator to include:</li> </ul>
734		Settings and monitored data
735		<ul> <li>Graphics-waveforms and loops (if available)</li> </ul>
736		• An evaluation of the patient's response to ventilation support (to include but not limited
737		to):
738		<ul> <li>Breath sounds, vital signs, and physical appearance</li> </ul>
739		<ul> <li>Arterial blood gases (if available)</li> </ul>
740		<ul> <li>Data from non-invasive monitors, e.g. SpO2 and ETCO2</li> </ul>
741		<ul> <li>Chest radiograph (if available)</li> </ul>
742		
743		
744	Action to	be taken in the event of an acute deterioration in the patient's clinical condition.
745		be taken in the event of an acute deterioration in the patient 3 clinical condition.
746	In the eve	ent of an acute deterioration in the patient's condition during the course of mechanical
746 747		ent of an acute deterioration in the patient's condition during the course of mechanical $a_{1} = a_{2} + a_{3} + a_{4} + a_{5} + a_{$
747	ventilatior	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop
747 748	ventilation of > 20 mm	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory
747 748 749	ventilation of > 20 mm	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop
747 748 749 750	ventilatior of > 20 mn care pract	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will
747 748 749 750 751	ventilatior of > 20 mn care pract	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will
747 748 749 750 751 752	ventilation of > 20 mm care pract	h as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will nmediately notify the nurse and provider. lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.
747 748 749 750 751 752 753	ventilation of > 20 mm care pract	h as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will nmediately notify the nurse and provider. lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0. ssess the patient to rule out one of the following conditions:
747 748 749 750 751 752 753 754	ventilation of > 20 mm care pract	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will nmediately notify the nurse and provider. lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0. ssess the patient to rule out one of the following conditions: o Acute airway obstruction
747 748 749 750 751 752 753	ventilation of > 20 mm care pract	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will nmediately notify the nurse and provider. lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0. ssess the patient to rule out one of the following conditions: • Acute airway obstruction
747 748 749 750 751 752 753 754 755	ventilation of > 20 mm care pract	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will nmediately notify the nurse and provider. lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0. ssess the patient to rule out one of the following conditions:
747 748 749 750 751 752 753 754 755 756	ventilation of > 20 mm care pract	n as evidenced by acute oxygen desaturation (SpO2 < 80%), acute hypotension (mean BP drop mHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will nmediately notify the nurse and provider. lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0. ssess the patient to rule out one of the following conditions:
747 748 749 750 751 752 753 754 755 756 757 758 759	ventilation of > 20 mm care pract	<ul> <li>as evidenced by acute oxygen desaturation (SpO2 &lt; 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will</li> <li>nmediately notify the nurse and provider.</li> <li>lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.</li> <li>ssess the patient to rule out one of the following conditions: <ul> <li>Acute airway obstruction</li> <li>Bronchospasm</li> <li>Pneumothorax</li> <li>Flash pulmonary edema</li> <li>Aspiration</li> <li>Airway misplacement – e.g. accidental extubation or decannulation, intubation of the</li> </ul> </li> </ul>
747 748 749 750 751 752 753 754 755 756 757 758 759 760	ventilation of > 20 mm care pract	<ul> <li>as evidenced by acute oxygen desaturation (SpO2 &lt; 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will</li> <li>nmediately notify the nurse and provider.</li> <li>lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.</li> <li>ssess the patient to rule out one of the following conditions: <ul> <li>Acute airway obstruction</li> <li>Bronchospasm</li> <li>Pneumothorax</li> <li>Flash pulmonary edema</li> <li>Aspiration</li> <li>Airway misplacement – e.g. accidental extubation or decannulation, intubation of the right-mainstem bronchus.</li> </ul> </li> </ul>
747 748 749 750 751 752 753 754 755 756 757 758 759 760 761	ventilation of > 20 mm care pract	<ul> <li>as evidenced by acute oxygen desaturation (SpO2 &lt; 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will</li> <li>nmediately notify the nurse and provider.</li> <li>lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.</li> <li>ssess the patient to rule out one of the following conditions: <ul> <li>Acute airway obstruction</li> <li>Bronchospasm</li> <li>Pneumothorax</li> <li>Flash pulmonary edema</li> <li>Aspiration</li> <li>Airway misplacement – e.g. accidental extubation or decannulation, intubation of the right-mainstem bronchus.</li> <li>Equipmentfailure</li> </ul> </li> </ul>
747 748 749 750 751 752 753 754 755 756 757 758 759 760 761 762	ventilation of > 20 mm care pract	<ul> <li>as evidenced by acute oxygen desaturation (SpO2 &lt; 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will</li> <li>nmediately notify the nurse and provider.</li> <li>lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.</li> <li>ssess the patient to rule out one of the following conditions: <ul> <li>Acute airway obstruction</li> <li>Bronchospasm</li> <li>Pneumothorax</li> <li>Flash pulmonary edema</li> <li>Aspiration</li> <li>Airway misplacement – e.g. accidental extubation or decannulation, intubation of the right-mainstem bronchus.</li> <li>Equipmentfailure</li> <li>System leak/disconnect</li> </ul> </li> </ul>
747 748 749 750 751 752 753 754 755 756 757 758 759 760 761 762 763	ventilation of > 20 mm care pract	<ul> <li>as evidenced by acute oxygen desaturation (SpO2 &lt; 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will</li> <li>nmediately notify the nurse and provider.</li> <li>lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.</li> <li>ssess the patient to rule out one of the following conditions: <ul> <li>Acute airway obstruction</li> <li>Bronchospasm</li> <li>Pneumothorax</li> <li>Flash pulmonary edema</li> <li>Aspiration</li> <li>Airway misplacement – e.g. accidental extubation or decannulation, intubation of the right-mainstem bronchus.</li> <li>Equipmentfailure</li> <li>System leak/disconnect</li> </ul> </li> </ul>
747 748 749 750 751 752 753 754 755 756 757 758 759 760 761 762 763 764	ventilation of > 20 mm care pract	<ul> <li>as evidenced by acute oxygen desaturation (SpO2 &lt; 80%), acute hypotension (mean BP drop mHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will</li> <li>nmediately notify the nurse and provider.</li> <li>lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.</li> <li>ssess the patient to rule out one of the following conditions: <ul> <li>Acute airway obstruction</li> <li>Bronchospasm</li> <li>Pneumothorax</li> <li>Flash pulmonary edema</li> <li>Aspiration</li> <li>Airway misplacement – e.g. accidental extubation or decannulation, intubation of the right-mainstem bronchus.</li> <li>Equipment failure</li> <li>System leak/disconnect</li> </ul> </li> </ul>
747 748 749 750 751 752 753 754 755 756 757 758 759 760 761 762 763	ventilation of > 20 mm care pract	<ul> <li>as evidenced by acute oxygen desaturation (SpO2 &lt; 80%), acute hypotension (mean BP drop nHg), an acute increase in airway pressure or an acute decrease in tidal volume the respiratory titioner will</li> <li>nmediately notify the nurse and provider.</li> <li>lanually ventilate the patient with a manual resuscitator set to deliver a FiO2 of 1.0.</li> <li>ssess the patient to rule out one of the following conditions: <ul> <li>Acute airway obstruction</li> <li>Bronchospasm</li> <li>Pneumothorax</li> <li>Flash pulmonary edema</li> <li>Aspiration</li> <li>Airway misplacement – e.g. accidental extubation or decannulation, intubation of the right-mainstem bronchus.</li> <li>Equipmentfailure</li> <li>System leak/disconnect</li> </ul> </li> </ul>

- 766 Appendix A: FiO2/PEEP Table
- 767 768
- 769 Oxygenation Goals:
- 770 771

774

779 780

787

792 793

794

- $60 \,\mathrm{mm}\,\mathrm{Hg} \le \mathrm{PaO2} \le 85 \,\mathrm{mm}\,\mathrm{Hg}$  (
- 773 □ SaO2 > 90%
  - SpO2 > 90%
- 775 776 Clinical Application of Standard: If not at goal, move up one step, if above goal(s) move down one step -PEEP adjustments within each step are based on clinical assessment. 777 778
  - PEEP FiO2 <.40 5 0.40-0.60 5-8 > 0.60 8-15
- 786 **Appendix B: Arterial Blood Gases**
- 788 An arterial blood gas sample should be obtained within 30 minutes following the initiation of mechanical 789 ventilation.
- 790 791 Subsequent arterial blood gas samples should be obtained...
  - Upon receipt of a provider's order
  - □ Following a ventilator setting change that is intended to stabilize or achieve ventilation and/or acid-base goals.
- 796 Following a ventilator change that is intended to stabilize or achieve oxygenation goals for the patient receiving total mechanical ventilation support (when non-invasive monitoring is 797 unavailable to insufficient to provide reliable data). 798
- 799 To assess the oxygenation, ventilation, and acid-base effects resulting from the changeover to 800 partial mechanical ventilation support from total mechanical ventilation support.
- 801 To assess the oxygenation, ventilation, and acid-base effects resulting from a ventilator setting change that leads to a significant change in respiratory rate, tidal volume and/or minute volume 802 803 for patients in either total or partial mechanical ventilation support.
- 804
- 805
- 806 **Appendix C: Tolerance Criteria**

807 808	Patien exists:	t may be considered intolerant of the partial support settings or a SBT if any of the following
808 809	exists.	
810 811		Development of rapid, shallow breathing: $f \ge 35$ or an increase of $\ge 10$ breaths per minute over previous respiratory rate.
812 813		Intolerable dyspnea, diaphoresis, excessive use of accessory muscles, or development of paradoxical respirations.
814 815 816 817 818		Heart rate > 120 or a change in heart rate of $\geq$ 20 that cannot be attributed to another cause. Diastolic blood pressure change of 20 mm Hg that cannot be attributed to another cause. Development of cardiac arrhythmia, deterioration of mental state, or deterioration of arterial blood gases.
819	Appen	dix D. Daily Spontaneous Breathing Trial
820 821 822 823 824 825	extuba	atient requiring mechanical ventilation should be assessed daily to determine readiness for tion and discontinuation of mechanical ventilation support with a Spontaneous Breathing Trial In patients requiring mechanical ventilation for more than 21 days, SBTs should be considered n PSV.
826 827 828 829 830 831 832 833		<ul> <li>The RCP will conduct a SBT on patients who meet the following indications:</li> <li>FiO2 ≤0.4</li> <li>PEEP ≤8 cm H2O</li> <li>A reliable, spontaneous respiratory drive regardless of mode or level of support provided.</li> <li>Tolerance criteria outlined in Appendix C is met.</li> <li>The patient's overall condition is stable or improving.</li> </ul>
834 835 836 837 838 839 840 841		<ul> <li>receiving neuromuscular blockade</li> <li>on HFOV</li> <li>with an inspiratory to expiratory ratio ≥ 1:1</li> <li>with a pH ≤ 7.20</li> <li>with an impending MI</li> <li>with a BP systolic of &lt; 80; MAP &lt; 60, or HR &gt; 120 and/or the need for vasopressor therapy (dopamine or dobutamine) ≥ 10 ∞g/kg/min, or more than one vasopressor b maintain hemodynamic stability</li> </ul>
842 843		The SBT will be performed with a low level of CPAP (5-8 cm H2O), PSV 5 with 5-8 cm H2O PEEP, or automatic tube/airway compensation (ATC/AAC) while maintaining the ventilator's FiO2.
844		The head of the bed should be elevated to 30 degrees.
845 846 847 848		The RCP will monitor the patient closely for the first five minutes of the SBT to assess tolerance and, if tolerated the SBT will continue for at least 30 minutes but not greater than 120 minutes. Patients who tolerate the SBT will be considered for extubation (see Tolerance Criteria in Appendix C).
849 850 851 852 853		At the completion of the trial, the following will be documented in the medical record: <ul> <li>Mode (e.g., PS-SBT, CPAP-SBT or ATC/AAC)</li> <li>Respiratory rate</li> <li>Minute volume</li> <li>Tidal volume</li> </ul>

854		∘ f/VT
855 856		Patients who fail the SBT trial will be returned to mechanical ventilation at previously tolerated settings.
857		A note will be entered into the medical record to indicate the reason for failure (e.g., high f/VT).
858 859		A note will be entered into the medical record to indicate the reason a SBT was not indicated in patients not receiving an SBT despite meeting criteria above
860		Extubation
861 862		<ul> <li>The RCP will recommend extubation or discontinuing mechanical ventilation support when the patient meets the extubation criteria listed in Appendix F.</li> </ul>
863		$\circ$ The RCP will extubate the patient upon receipt of a provider's order.
864 865	۸nnon	dix E. Spontaneous Breathing Trials in tracheotomized patients
866	Abben	aix L. Spontaneous Dreathing mais in tracheotomized patients
867		Spontaneous breathing trials for tracheotomized patients combine periods of spontaneous
868 869		breathing, generally of increasing duration, with periods of mechanical ventilation support for patients who meet criteria in Appendix D.
870		The decision to initiate and conduct these trials will be made by the medical care team as part of
871		the daily plan of care. A provider's order is required to initiate a spontaneous breathing trial for
872		tracheotomized patients.
873 874		The patient will be removed from mechanical ventilation support and placed on a High Flow device. Flow and FiO2 will be adjusted to meet SpO2 goal.
875		The patient will be placed on a continuous end tidal CO2 monitor with alarms set $10$ mmHG
876		above and below their established baseline end tidal CO2.
877 878		• *ETCO2 will not be monitored during Passey-Muir Valve use.
878 879		<ul> <li>*ETCO2 may be discontinued once the patient is successfully liberated from mechanical ventilation &gt; 48 hours.</li> </ul>
880		Patient tolerance will be assessed (Appendix C).
881		The RCP will return the patient to mechanical ventilation support
882		<ul> <li>If the patient fails to meet tolerance criteria, or</li> </ul>
883		<ul> <li>According to the time interval determined by the medical team (plan of care), or</li> </ul>
884		<ul> <li>To "rest" the patient overnight with the intention of continuing trial the following</li> </ul>
885		morning.
886	Annon	dix E. Evilubation Critaria
887 888	Appen	dix F: Extubation Criteria
889 890	A patie	ent should be considered for extubation when the following criteria are met:
891		Patient is able to tolerate a Spontaneous Breathing Trial.
892		Adequate airway protection, a reliable respiratory drive, and airway suctioning no more
893		frequently than every two hours.
894		${\tt Successful}\ `{\tt cuff} {\tt leak}\ {\tt test}\ `{\tt in}\ {\tt patients}\ {\tt suspected}\ {\tt of}\ {\tt possible}\ {\tt upper}\ {\tt airway}\ {\tt abnormalities}.$
895		NIFM ≥ -25 (Lung Transplant Patients)
896		Diaphragm unclamped (Lung Transplant Patients)
897		
898		

**REFERENCES** 

900 Epoprostenol: Adult Inhaled Pulmonary Vasodilator Protocol

### 903 904 PATIENT POPULATION:

Patients receiving Nitric Oxide (iNO) for pulmonary vasodilator therapy are eligible for continuous
 aerosolized or inhaled epoprostenol .

# 909 **DESCRIPTION**:

901 902

905

908

917

920

- 910
  911 Veletri is a naturally occurring prostaglandin that serves as a potent vasodilator and is an effective
  912 inhibitor of platelet aggregation. Aerosolized Veletri is used as a selective pulmonary vasodilator when
  913 administered by inhalation, it has been shown to improve oxygenation, reduce pulmonary shunt,
- 914 lower pulmonary artery pressure and pulmonary vascular resistance.

### 915 916 **INDICATIONS**:

918 Post-cardiothoracic surgery patients located on 7 West for management of pulmonary hypertension, 919 right ventricular dysfunction, or refractory hypoxemia.

## 921 **PREPARATION/STORAGE/DISPENSING:**

- 922 Uveletri syringes shall be prepared by the pharmacy sterile preparation cleanroom (SPC). Standard concentration of syringe is 1.5 mg/50 mL (30,000 ng/mL).
- 924 D Prepared Veletri syringes shall be given a 7 days refrigerated beyond use dating.
- A non-patient specific supply of syringes will be stored in the Omnicell controlled refrigerator in the "B"
   medication room on 7W. Only Respiratory Medications will be stored in this refrigerator.

927	
928	To assure ongoing appropriate inventory management, respiratory therapists shall access the
929	Veletri syringes by logging into the Omnicell cabinet and selecting the drug for removal each
930	time.
931	
932	ADMINISTRATION GUIDELINES:
933	
934	1. Patient will present with Veletri from the operation room at a dosage rate of 50
935	ng/kg/min.
936	2. The usual dosage range is 15-50 ng/kg/min based on ideal body weight (IBW).
937	3. The usual starting dose of aerosolized Veletri is 50 ng/kg/min.
938	4. The dose of 50ng/kg/min is at the upper end of the dose range used in most clinical
939	studies.
940	5. In order to increase the likelihood of a timely response it is preferable to start at 50
941	ng/kg/min and then, if needed, titrate dose down once a favorable response has been
942	confirmed and the patient is stable.
942 943	6. Dosage adjustments should be made in increments/decrements of 5ng/kg/min.
943 944	
	7. Any variation from the standard concentration for <b>inhaled use</b> will require approval
945 046	from RT and pharmacy administration.
946	8. Syringe must be changed at a minimum of every 24 hours; nebulizer and connecting
947	tubing changed every 7 days.
948	9. Veletri is administered using a special nebulizer set-up as described below.
949	10. Veletri can be administered during invasive or noninvasive ventilation, and via facemask
950	orHighFlownasalcannula.
951	
951 952	ASSESSMENT OF RESPONSE:
951 952 953	ASSESSMENT OF RESPONSE:
951 952 953 954	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved
951 952 953 954 955	ASSESSMENT OF RESPONSE:
951 952 953 954 955 956	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.
951 952 953 954 955 956 957	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved
951 952 953 954 955 956 957 958	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation. PRECAUTIONS AND SIDE EFFECTS:
951 952 953 954 955 956 957 958 959	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation. PRECAUTIONS AND SIDE EFFECTS: 1. Inhaled Veletri has fewer adverse effects than intravenous administration.
951 952 953 954 955 956 957 958 959 960	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation. PRECAUTIONS AND SIDE EFFECTS: 1. Inhaled Veletri has fewer adverse effects than intravenous administration. 2. Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare
951 952 953 954 955 956 957 958 959 960 961	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation. PRECAUTIONS AND SIDE EFFECTS: 1. Inhaled Veletri has fewer adverse effects than intravenous administration. 2. Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug,
951 952 953 954 955 956 957 958 959 960 961 962	ASSESSMENT OF RESPONSE: The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation. PRECAUTIONS AND SIDE EFFECTS: 1. Inhaled Veletri has fewer adverse effects than intravenous administration. 2. Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.
951 952 953 954 955 956 957 958 959 960 961 962 963	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> <li>Abrupt withdrawal of inhaled Veletri can cause rebound pulmonary vasoconstriction</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> <li>Abrupt withdrawal of inhaled Veletri can cause rebound pulmonary vasoconstriction and hypoxemia, but this is rare.</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 967	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> <li>Abrupt withdrawal of inhaled Veletri can cause rebound pulmonary vasoconstriction</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 967 968	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> </ol> </li> <li>EQUIPMENT:</li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 967 968 969	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> </ol> </li> <li>EQUIPMENT: <ol> <li>Medfusion 3500 Infusion pump</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 965 966 967 968 969 969 970	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> <li>Abrupt withdrawal of inhaled Veletri can cause rebound pulmonary vasoconstriction and hypoxemia, but this is rare.</li> </ol> </li> <li>EQUIPMENT: <ol> <li>Medfusion 3500 Infusion pump</li> <li>Aeroneb Pro-xcontrol unit, Aeroneb Pro-x nebulizer with T adapter, Aerogen Tubing Set</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 967 966 967 968 969 970 970	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> <li>Abrupt withdrawal of inhaled Veletri can cause rebound pulmonary vasoconstriction and hypoxemia, but this is rare.</li> </ol> </li> <li>EQUIPMENT: <ol> <li>Medfusion 3500 Infusion pump</li> <li>Aeroneb Pro-xcontrol unit, Aeroneb Pro-x nebulizer with T adapter, Aerogen Tubing Set</li> <li>50mL Veletri syringe prepared by pharmacy</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 967 968 967 968 969 970 971 972	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> <li>Abrupt withdrawal of inhaled Veletri can cause rebound pulmonary vasoconstriction and hypoxemia, but this is rare.</li> </ol> </li> <li>EQUIPMENT: <ol> <li>Medfusion 3500 Infusion pump</li> <li>Aeroneb Pro-xcontrol unit, Aeroneb Pro-x nebulizer with T adapter, Aerogen Tubing Set</li> <li>50 mL Veletri syringe prepared by pharmacy</li> <li>Hydroscopic filters to be placed between expiratory limb and ventilator exhalation filter)</li> </ol> </li> </ul>
951 952 953 954 955 956 957 958 959 960 961 962 963 964 965 966 967 966 967 968 969 970 970	<ul> <li>ASSESSMENT OF RESPONSE:</li> <li>The desired response to inhaled Veletri is a decrease in pulmonary artery pressure, improved hemodynamics, and/or improved arterial oxygenation.</li> <li>PRECAUTIONS AND SIDE EFFECTS: <ol> <li>Inhaled Veletri has fewer adverse effects than intravenous administration.</li> <li>Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except at very high doses. If hypotension should occur when initiated, stop the drug, reinitiate iNO and contact the ICU team.</li> <li>Veletri has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.</li> <li>Abrupt withdrawal of inhaled Veletri can cause rebound pulmonary vasoconstriction and hypoxemia, but this is rare.</li> </ol> </li> <li>EQUIPMENT: <ol> <li>Medfusion 3500 Infusion pump</li> <li>Aeroneb Pro-xcontrol unit, Aeroneb Pro-x nebulizer with T adapter, Aerogen Tubing Set</li> <li>50mL Veletri syringe prepared by pharmacy</li> </ol> </li> </ul>

97<del>6</del>



# 978 **PRACTITIONER ROLES:** 979

010		
980	Nursing/Respiratory T	herapy Considerations
981		
982	1.	Any questions about the dose should be clarified with the pharmacist.
983	2.	This therapy demands close collaboration between the respiratory therapist,
984 085	0	nurse, and pharmacist.
985	3.	Drug administration is documented in E-MAR, Respiratory Medication
986		Administration flowsheet and noted in the shift note by the respiratory
987	4	therapist.
988 989	4. 5	Continuous oxygen saturation monitoring is required.
969 990	5.	Standard unit-specific vital signs assessment is followed.
990 991	6. 7.	Aerosol delivery into the ventilator circuit or mask must be confirmed visually. If the patient is receiving Veletri by face mask, assure that the mask is fitted
991 992	1.	comfortably and is only removed for short periods; monitor oxygen saturation
992 993		closely when the mask is removed.
995 994	8.	Care should be taken to avoid direct exposure to the aerosol emitted from the
995	0.	nebulizer.
996	9.	Masks are not required when entering the room or when involved in usual
997	0.	caregiver activities in the room. However, N95 masks will be available for those
998		who choose to use them.
999	10.	Although evidence is lacking for exposure during pregnancy, it is recommended
1000		that women who are pregnant do not enter the room during treatment.
1001		
1002	EQUIPMENT SET-UP:	
1003		
1004	1.	Assure Aeroneb control unit is plugged into Uninterruptible Power Source (UPS)
1005	2.	Assure that UPS and infusion pump are plugged into 110 volt AC power source.
1006	3.	Place additional expiratory filter between expiratory limb and ventilator
1007		expiratory filter.(Q4H and prn change)
1008	4.	Confirm the dose in the E-MAR. A dose greater than 50 ng/kg/min will not be
1009		used. If there are concerns about spillover systemic hypotension, a lower
1010		starting dose (e.g., 30 ng/kg/min) should be used.
1011	5.	Scan the patient's ID band, scan barcode on syringe label, and look for Maestro
1012		Medication Administration window to appear.
1013	6.	Perform functional check of nebulizer.
1014		<ul> <li>Inject 1 mL of sterile saline into nebulizer cup and t-piece assembly.</li> </ul>
1015		<ul> <li>Press and hold the On/Off power button for 3 seconds.</li> </ul>
1016		
1017		
1018		page 22
		r-8

- Verify that Continuous Mode indicator illuminates and aerosol is produced.
- Allow nebulizer cup to empty.
- 7. Attach Aerogen tubing set to syringe containing Veletri solution. Attach the other end of the tubing to the luer connector for the nebulizer; this connector should be separate from the nebulizer assembly at this time.



# Program Medfusion 3500 Infusion pump.

• Power pump on

) _0	- CHENNENH				CID
	In Priling Street	E Plepa	CATTO	h Sar	CTV C
					Calles 0
	C				- 201 - 6

□ Select Respiratory Folder



Select Epoprostenol Folder

#### 



 $\Box$  Verify that the concentration printed on the syringe label is 1.5 mg/50mL and press "Yes".

1039	
1849	
1042 1043 1044	<ul> <li>Choose B-D syringe.</li> <li>Place 60 mL syringe containing 50 mL Epoprostenol (Veletri) solution into infusion pump and press "Enter".</li> </ul>
1045 1046	Enter patient's ideal body weight (ideal body weight) in KG
1047 1048 1049	□ Enter starting dose 50 ng/kg/min as ordered
1050 1051 1052 1053 1054	<ul> <li>Prime the tubing by press and holding BOLUS until tubing is primed, (approximate priming volume 3.7 ccs')</li> <li>Confirm no "air" is in Aerogen tubing</li> </ul>
1055 1056 1057	<ul> <li>The dose set on the pump will be confirmed by a second clinician (respiratory therapist or nurse).</li> <li>Attach tubing from Veletri syringe to pebulizer.</li> </ul>

- 1057 Description Attach tubing from Veletri syringe to nebulizer.
- 1058 Delace Aeroneb nebulizer unit into T-piece.
- 1059 Determination of the termination of term
- 1060Press START (green button) on pump to begin dose after nebulizer is in circuit. Observe1061nebulizer for aerosol production.

	a dia	The second secon
	and the second	
1063	) 🥥	
1863		
1065		Response to therapy should be assessed after 30-60 minutes. Determination of criteria for
1066		response should be established by the clinical team before therapy is initiated. A general
1067		guideline is that there should be a 20% improvement in oxygenation or hemodynamics to
1068		continue therapy after 1 hour.
1069 1070		Maximum dose is 50ng/kg/min. A lack of response at this dose level should prompt discontinuation of therapy
1071		If hypotension occurs, dose should be reduced.
1072		Dosage adjustments should be made in increments/decrements of 5 ng/kg/min.
1073		After 4 hours of clinical stability, consideration should be given to dose reduction.
1074		
1075	Changin	g Dose:
1076	_	
1077		Select Change dose (ng/kg/min)
1078 1079		Enter new dose, press start.
1079	Syringe	Change:
1081	Oyninge	onange.
1082		The Veletri syringe is changed at least every 24 hrs or whenever the remaining volume in the
1083		syringe reaches 5 mL
1084		The connecting tubing and nebulizer only need to be changed every 7 days.
1085		$\circ$ (The syringe will need to be changed in less than 24 hrs when the infusion rate is greater
1086		than 1.8 mL/hr).
1087		
1088 1089		<ol> <li>To change syringe press STOP</li> <li>Remove tubing from nebulizer and close</li> </ol>
1089		<ol> <li>Remove tubing from nebulizer and close port</li> </ol>
1090		3. Remove syringe
1092		4. Remove tubing from syringe and attach to
1093		new syringe
1094		5. Place new syringe in pump and Prime
1095		tubing
1096		6. Reattach tubing to nebulizer
1097		7. Press START
1098		8. Oberve for delivery and aerosol
1099 1100		production
1100		

-----

#### 1101 Invasive ventilator assembly: 1102

1103

- 1. Connect T-piece into circuit at humidifer inlet (dry side); be certain that nebulizer cup is upright.
- 1104 2. Place disposable bacterial/viral filter before exhalation valve assembly. 1105
  - Change the filter every 4 hrs and PRN if resistance to expiratory flow is noted.

 3. An active humidification system must be used. Do not use a Heat-Moisture Exchanger (HME) during administration of inhaled Veletri.



# 1111 Transport Ventilator Assembly:

- 1. Place filter on ventilator outlet port
- 2. Connect T-piece/aerogen assembly into filter outlet.
- 3. Insert Expiratory filter on limb
  - 4. Be certain that nebulizer cup is upright.
    - 5. Do not use a Heat-Moisture Exchanger (HME) during administration of inhaled Veletri.



# 

- 1. Use a non-vented mask.
- 2. Place the nebulizer between the leak port and the mask.
- 3. Be careful to position the mask so that the nebulizer cup remains in a vertical position.



Noninvasive Ventilator Assembly:

## 1126 Mask Assembly:

I

- 1. Attach Aerogen "Mask Adapter" to Aerosol Mask inlet

1133

1134

- 2. Connect small bore O2 tubing to bottom inlet port of adapter
  - 3. Place Aerogen nebulizer in Aerogen adapter



#### 1135 High flow Nasal Cannula Assembly: 1136

- 1. Connect T piece to humidifier inlet with the nebulizer cup in an upright, vertical position.
- 2. Connect high flow nasal cannula to the other end of T piece.
- 3. Adjust oxygen flow meter to desired flow rate (liters/minute)

#### 1139 1140

1137

1138



# 1141

# Manual Ventilation Assembly:

- 1. For patients with an artificial airway, place the T piece between the manual ventilator (AMBU) outlet and the endotracheal or tracheostomy tube. Be certain that the nebulizer cup is in a vertical position.
- 2. If a non-intubated patient requires emergency ventilation, immediately initiate manual bagvalve-mask ventilation.
- 3. If emergency ventilation is needed, the first priority is adequate ventilation and oxygenation and the secondary priority is administration of aerosolized Veletri.



# **Equipment Maintenance**

1153 1154 1155

1152

1. Disposable hydroscopic filters are to be changed <u>Q4h and prn</u>.

1142

1143 1144

1145

1146 1147

1148 1149

1156	
1157	2. Aeroneb nebulizer to be changed Q7 days and prn.
1158	3. Syringe line tubing to be changed Q7 days and prn.
1159	
1160	RECOMMENDED DOSING STRATEGIES:
1161	
1162	1. The usual dosage range is 5-50 ng/kg/min based on ideal body weight. Do not use a dose greater
1163 1164	than 50ng/kg/min.
1164 1165	<ol> <li>The usual starting dose of aerosolized Veletri is 50ng/kg/min.</li> <li>Dosage adjustments should be made in increments/decrements of 5ng/kg/min.</li> </ol>
1166	<ol> <li>The initial dose of 50ng/kg/min is at the upper end of the dose range used in most clinical</li> </ol>
1167	studies.
1168	5. Assess the response to therapy within 30-60 minutes of initiation.
1169	6. After 4 hours of clinical stability, consideration should be given to dose reduction.
1170	7. At high doses, there is a potential for systemic effects, which results in systemic hypotension.
1171	There is a potential for rebound pulmonary vasoconstriction and hypoxemia when Veletri is
1172	abruptly discontinued, but it is probably less than that for inhaled nitric oxide.
1173	8. Be prepared to increase the FIO2 and support hemodynamics during discontinuation. If rebound
1174	occurs, it may be helpful to wean the dose slowly before discontinuation.
1175	
1176	
1177	
1178	
1179	WEANING
1180	
1181	GENERAL COMMENTS: Right Heart hemodynamics include: SVO2, CO, CI, PA sys/dia, CVp. There is a
1182	
	potential for rebound increase in pulmonary vasoconstriction and hypoxemia when Veletri is
1183	abruptly discontinued. Weaning of Epoprostenol will commence following communication and
1184	consent of the Provider team. Prior to each dosing change, Right Heart hemodynamics will be
1185	assessed and documented in the comment section of the MAR for Epoprostenol by inhalation. In
1186	general, the Right Heart parameters should meet the following criteria before weaning or
1187	discontinuation: SvO2>65, CVP < 15,="" ci="">2.2, and adequate oxygenation. Pulmonary arterial
1188	pressure values will vary but should remain stable during weaning and trial off.

## 

Dose	Action
50	Dosage rate at start of wean
45	Observe Right Heart hemodynamics and decrease dose to
	45ng/kg/min only and allow stabilizing for 1 hour.
35	Observe Right Heart hemodynamics and decrease dose to
30	35ng/kg/min only and allow stabilizing for 1 hour.
25	Observe Right Heart hemodynamics and decrease dose to
25	25ng/kg/min only and allow stabilizing for 1 hour.
15	Observe Right Heart hemodynamics and decrease dose to
	15ng/kg/min only and allow stabilizing for 1 hour.
	If Right Heart hemodynamics are stable, notify Provider for
Off	Discontinue of Epoprostenol and observe for 1 hour before taking
	down setup

  If rebound worsening of pulmonary hypertension, reduced SvO2 < 65, increased cvp by more than 5 mm Hg or cvp value is > 15 mm Hg, reduced CI < 2.2, or hypoxemia occurs during any dosing decrease, return to last dose, and notify provider team. Provider may choose to resume wean in 1 hour or halt weaning at

current dose until further discussion.

# DISCONTINUATION:

WEANING PROCEDURE:

1. Once weaning protocol has been completed, notify the provider team of hemodynamic stability with Epoprostenol off after 1 hour.

1. Obtain consent for weaning commencement from provider team.

2. The following decrement strategy will be used:

2. If rebound worsening of pulmonary hypertension, reduced SvO2 < 65, increased cvp by more than 5 mm Hg or cvp value is > 15 mm Hg, reduced CI < 2.2, or hypoxemia occurs during any dosing decrease, return to last dose, and notify provider team. Provider may choose to resume wean in 1 hour or halt weaning at current dose until further discussion.</p>

#### **DOCUMENTATION**

- 1. MAR documentation: Scan patient and scan medication. Enter dose and rate as delivered.
- 2. Respiratory Medication administration Documet continuous nebulizer on Respiratory Medication Administration flowsheet. Use this comment section for care notes.

# CAREGIVER PROTECTION CONCERNS:

1. Toxicity: Veletri has no known toxic effects or toxic metabolites.

1225	2. Carcinogenesis, Mutagenesis, Impairment of Fertility: L	ong-term studies in animals have not
1226	been performed to evaluate carcinogenic potential. A mic	cronucleus test in rats revealed no
1227	evidence of mutagenicity. The Ames test and DNA elution t	ests were also negative, although the
1228	instability of Veletri makes the significance of these tests u	•
1229	rats given Veletri by subcutaneous injection at doses up to	-
1230	dose.	
1231	<ol> <li>Pregnancy: Pregnancy Category B. Reproductive studies h</li> </ol>	ave been performed in pregnant rats
1232	and rabbits at doses up to 2.5 times the recommended h	
1232	evidence of impaired fertility or harm to the fetus due to	
1233	adequate and well-controlled studies in pregnant women.	
1234	4. Recommendations:	
1235		it do as not pood to be a popotiva
1236	• The patient should be in a single patient room, but	•
1237	pr <b>essure room</b> and the door does not need to remain	
1230	<ul> <li>A sign will be placed on the door to indicate that <i>inha</i></li> <li>Care should be taken to avoid direct exposure to the factor of the statement of the statemen</li></ul>	•
1239	<ul> <li>Care should be taken to avoid direct exposure to the nebulizer.</li> </ul>	le delosol enlitted from the
1240		when involved in usual coregiver
1241	<ul> <li>Masks are not required when entering the room or activities in the room, but N95 masks will be availa</li> </ul>	•
1242	them.	ole for those who choose to use
1243	<ul> <li>Women who are pregnant should not enter the row</li> </ul>	am during traatmont
1244	o women who are pregnant should not enter the to	on during treatment.
1246		
1247	Storage and Cleaning of System	
1248		
1249	1. Disposable equipment, (nebulizer, t-piece, syringe, syringe tubin	g) will be removed and discarded in the
1250	patient room upon termination of use.	-
1250 1251	<ul><li>patient room upon termination of use.</li><li>2. The administration system will be disinfected with "Sani-wipes"</li></ul>	-
1250 1251 1252	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> </ul>	prior to removal from the patients' room
1250 1251 1252 1253	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets who have a structure of the storage room for the storage room f</li></ul>	prior to removal from the patients' room
1250 1251 1252 1253 1254	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> </ul>	prior to removal from the patients' room
1250 1251 1252 1253 1254 1255	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets wh charge.</li> </ol>	prior to removal from the patients' room
1250 1251 1252 1253 1254 1255 1256	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets who have a structure of the storage room for the storage room f</li></ul>	prior to removal from the patients' room
1250 1251 1252 1253 1254 1255 1256 1257	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets who charge.</li> </ul>	prior to removal from the patients' room
1250 1251 1252 1253 1254 1255 1256 1257 1258	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets who charge.</li> <li><b>REFERENCES:</b></li> <li>1. Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bost</li> </ul>	orior to removal from the patients' room ile in storage to maintain full battery s AP. Randomized controlled trial of
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets where charge.</li> <li><b>REFERENCES:</b></li> <li>1. Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bogaerosolized prostacyclin therapy in children with acute lution.</li> </ul>	orior to removal from the patients' room ile in storage to maintain full battery s AP. Randomized controlled trial of
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets who charge.</li> <li><b>REFERENCES:</b> <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Boa aerosolized prostacyclin therapy in children with acute lu 32(4):1055-60.</li> </ol> </li> </ul>	orior to removal from the patients' room lile in storage to maintain full battery SAP. Randomized controlled trial of ng injury. Crit Care Med 2004;
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets where charge.</li> <li><b>REFERENCES:</b> <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bogaerosolized prostacyclin therapy in children with acute lug 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prostacyclin</li> </ol> </li> </ul>	orior to removal from the patients' room ile in storage to maintain full battery s AP. Randomized controlled trial of ng injury. Crit Care Med 2004; tacyclin (PGI2) in acute respiratory
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets where charge.</li> <li><b>REFERENCES:</b> <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bogaerosolized prostacyclin therapy in children with acute lut 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prostacyclin therapy (pulmonary injury) and anticed prostacy of primary (pulmonary injury) and primary (pulmonary injury)</li> </ol> </li> </ul>	orior to removal from the patients' room ile in storage to maintain full battery s AP. Randomized controlled trial of ng injury. Crit Care Med 2004; tacyclin (PGI2) in acute respiratory of secondary (extrapulmonary injury)
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263	<ul> <li>patient room upon termination of use.</li> <li>2. The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>3. Administration systems will be plugged in to electrical outlets where charge.</li> <li><b>REFERENCES:</b> <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Boa aerosolized prostacyclin therapy in children with acute lut 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prostacyclin therapy (pulmonary injury) ar disease on gas exchange response. Crit Care Med 2001; 290</li> </ol> </li> </ul>	orior to removal from the patients' room ile in storage to maintain full battery s AP. Randomized controlled trial of ng injury. Crit Care Med 2004; tacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62.
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets which arge.</li> <li>REFERENCES:         <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bogaerosolized prostacyclin therapy in children with acute lut 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prostacyclin therapy (pulmonary injury) ard disease on gas exchange response. Crit Care Med 2001; 296</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. Crit Care Med</li> </ol> </li> </ol>	orior to removal from the patients' room alle in storage to maintain full battery s AP. Randomized controlled trial of ng injury. Crit Care Med 2004; tacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62. ed 2005; 33(3 Suppl):S188-95.
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264 1265	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets where charge.</li> <li><b>REFERENCES:</b> <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Boa aerosolized prostacyclin therapy in children with acute lut 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prost distress syndrome: impact of primary (pulmonary injury) ar disease on gas exchange response. Crit Care Med 2001; 29(</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. Crit Care Med 4. Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, Wat</li> </ol> </li> </ol>	orior to removal from the patients' room ille in storage to maintain full battery SAP. Randomized controlled trial of ng injury. Crit Care Med 2004; tacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62. ed 2005; 33(3 Suppl):S188-95. urick R, Hinder F, Booke M. Inhaled
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets wh charge.</li> <li>REFERENCES:         <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bogaerosolized prostacyclin therapy in children with acute lu 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prost distress syndrome: impact of primary (pulmonary injury) and disease on gas exchange response. Crit Care Med 2001; 290</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. Crit Care Med 4. Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, Wat prostaglandin E1 for treatment of acute lung injury in severe.</li> </ol> </li> </ol>	orior to removal from the patients' room ille in storage to maintain full battery SAP. Randomized controlled trial of ng injury. Crit Care Med 2004; tacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62. ed 2005; 33(3 Suppl):S188-95. urick R, Hinder F, Booke M. Inhaled
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266 1267	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets wh charge.</li> <li>REFERENCES:         <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bogaerosolized prostacyclin therapy in children with acute lu 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prost distress syndrome: impact of primary (pulmonary injury) ar disease on gas exchange response. Crit Care Med 2001; 290</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. Crit Care Med 2001; 290</li> <li>Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, Wat prostaglandin E1 for treatment of acute lung injury in seve Analg 1998; 86(4):753-8.</li> </ol> </li> </ol>	orior to removal from the patients' room hile in storage to maintain full battery SAP. Randomized controlled trial of ng injury. Crit Care Med 2004; tacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62. ed 2005; 33(3 Suppl):S188-95. urick R, Hinder F, Booke M. Inhaled re multiple organ failure. Anesth
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266 1267 1268	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets where charge.</li> <li>REFERENCES:         <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Boa aerosolized prostacyclin therapy in children with acute lug 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prost distress syndrome: impact of primary (pulmonary injury) ard disease on gas exchange response. Crit Care Med 2001; 29(</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. Crit Care Med 2001; 29(</li> <li>Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, War prostaglandin E1 for treatment of acute lung injury in seve Analg 1998; 86(4):753-8.</li> <li>Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, War prostaglandin E1 for treatment of acute lung injury in seve Analg 1998; 86(4):753-8.</li> </ol> </li> </ol>	orior to removal from the patients' room hile in storage to maintain full battery a AP. Randomized controlled trial of ng injury. Crit Care Med 2004; htacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62. ad 2005; 33(3 Suppl):S188-95. urick R, Hinder F, Booke M. Inhaled re multiple organ failure. Anesth urick R, Hinder F, Booke M. Inhaled
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266 1267 1268 1269	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets wh charge.</li> <li>REFERENCES:         <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Bos aerosolized prostacyclin therapy in children with acute lu 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prost distress syndrome: impact of primary (pulmonary injury) ar disease on gas exchange response. Crit Care Med 2001; 290</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. CritCare Med 9001; 290</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. Maximum for prostaglandin E1 for treatment of acute lung injury in seve Analg 1998; 86(4):753-8.</li> <li>Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, Wat prostaglandin E1 for treatment of acute lung injury in seve Analg 1998; 86(4):753-8.</li> </ol> </li> </ol>	orior to removal from the patients' room hile in storage to maintain full battery a AP. Randomized controlled trial of ng injury. Crit Care Med 2004; htacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62. ad 2005; 33(3 Suppl):S188-95. urick R, Hinder F, Booke M. Inhaled re multiple organ failure. Anesth urick R, Hinder F, Booke M. Inhaled
1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1262 1263 1264 1265 1266 1267 1268	<ol> <li>patient room upon termination of use.</li> <li>The administration system will be disinfected with "Sani-wipes" and placed in a designated RCS Equipment storage room for 7W.</li> <li>Administration systems will be plugged in to electrical outlets where charge.</li> <li>REFERENCES:         <ol> <li>Dahlem P, van Aalderen WM, de Neef M, Dijkgraaf MG, Boa aerosolized prostacyclin therapy in children with acute lug 32(4):1055-60.</li> <li>Domenighetti G, Stricker H, Waldispuehl B. Nebulized prost distress syndrome: impact of primary (pulmonary injury) ard disease on gas exchange response. Crit Care Med 2001; 29(</li> <li>Lowson SM. Inhaled alternatives to nitric oxide. Crit Care Med 2001; 29(</li> <li>Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, War prostaglandin E1 for treatment of acute lung injury in seve Analg 1998; 86(4):753-8.</li> <li>Meyer J, Theilmeier G, Van Aken H, Bone HG, Busse H, War prostaglandin E1 for treatment of acute lung injury in seve Analg 1998; 86(4):753-8.</li> </ol> </li> </ol>	orior to removal from the patients' room hile in storage to maintain full battery a AP. Randomized controlled trial of ng injury. Crit Care Med 2004; htacyclin (PGI2) in acute respiratory d secondary (extrapulmonary injury) 1):57-62. ad 2005; 33(3 Suppl):S188-95. urick R, Hinder F, Booke M. Inhaled re multiple organ failure. Anesth urick R, Hinder F, Booke M. Inhaled

1271	
1272	6. Olschewski H, Ghofrani HA, Walmrath D, Schermuly R, Temmesfeld-Wollbruck B, Grimminger F,
1273	Seeger W. Inhaled prostacyclin and iloprost in severe pulmonary hypertension secondary to lung
1274	fibrosis. Am J Respir Crit Care Med 1999; 160(2):600-7.
1275	$\label{eq:constraint} 7.  PutensenC, HormannC, KleinsasserA, Putensen-HimmerG. Cardiopulmonary effects of$
1276	aerosolized prostaglandin E1 and nitric oxide inhalation in patients with acute respiratory
1277	distress syndrome. Am J Respir Crit Care Med 1998; 157(6 Pt 1): 1743-7.
1278	8. Siobal M. Aerosolized prostacyclins. Respir Care 2004; 49(6):640-52.
1279	9. Siobal MS, Kallet RH, Pittet JF, Warnecke EL, Kraemer RW, Venkayya RV, Tang JF. Description and
1280	evaluation of a delivery system for a erosolized prostacy clin. Respir Care 2003; 48 (8): 742-53.
1281	10. van Heerden PV, Barden A, Michalopoulos N, Bulsara MK, Roberts BL. Dose-response to inhaled
1282	aerosolized prostacyclin for hypoxemia due to ARDS. Chest 2000; 117(3):819-27.
1283	11. Afshari A, Brok J, Moller AM, Wetterslev J. Aerosolized prostacyclin for acute lung injury (ALI)
1284	and acute respiratory distress syndrome (ARDS). Cochrane Database Syst Rev 2010; (8):
1285	CD007733.
1286	12. Siobal MS, Hess DR. Are inhaled vasodilators useful in acute lung injury and acute respiratory
1287	distress syndrome? Respir Care 2010;55(2): 144-57; discussion 157-61.
1288	13. Khan TA, Schnickel G, Ross D, et al. A prospective, randomized, crossover pilot study of inhaled
1289	nitric oxide versus inhaled prostacyclin in heart and lung transplant recipients. J Thorac
1290	Cardiovasc Surg 2009; 138(6): 1417-24.
1291	
1292	REFERENCES

1293 1294 1295 1296 1297 1298	Transport of the Mechanically Ventilated CardioThoracic Patient on inhaled Nitric Oxide or inhaled Epoprostenol
1299 1300 1301	This policy sets forth the standards for transporting the mechanically ventilated cardiothoracic patient on inhaled nitric oxide (iNO) or inhaled Epoprostenol (iEPO) within Duke University Hospital by respiratory care practitioners.
1302 1303	Patient transport includes patient preparation, movement to and from the ICU, a diagnostic suite, Operating room, and time spent at the destination.
1304	<ul> <li>For transport, a mechanical ventilator designed for transport and the Pulmonary Vasodilator</li></ul>
1305	system in use (iEpo, iNO), will be utilized.
1306	ETCO2 will be monitored during all transports of mechanically ventilated patients.
1307	<ul> <li>All mechanically ventilated patients will be accompanied during transit by a respiratory care</li></ul>
1308	practitioner (RCP).
1309	If the patient is scheduled to undergo a procedure in the operating suite, the RCP may return to
1310	his/her assigned area after verbal hand off with the Attending Anesthesiologist.
1311 1312 1313	Description
1314	Mechanically ventilated patients on inhaled pulmonary vasodilators are transported within the
1315	hospital for diagnostic or therapeutic procedures that cannot be performed in an in-patient unit.
1316	To insure patient safety, respiratory care practitioners will make every effort to provide and
1317	maintain an appropriate and constant level of ventilation, oxygenation, and pulmonary
1318	vasodilator delivery during transport.
1319	The RCP will assess the patient's airway, ventilation and oxygen status prior to the transport. If
1320	the patient is hemodynamically unstable or if difficulty maintaining the patient's ventilation or
1321	oxygenation status is anticipated during transport, the RCP will voice these concerns to the
1322	medical team.

1323		
1324	Indicat	ions
1325		
1326		Transport of all patients using Respiratory Care owned equipment must be accompanied by a
1327 1328		Respiratory Care Practitioner.
1329	Hazards	s and Complications
1330		·
1331		Hyperventilation during manual ventilation may lead to respiratory alkalosis, cardiac
1332		dysrhythmia, hypotension, and decreased cerebral perfusion.
1333		Loss of PEEP/CPAP may result in hypoxemia and decreased oxygen delivery.
1334 1335		Disruption of inhaled pulmonary vasodilator may cause rebound pulmonary hypertension and/or hypoxemia.
1336		Position changes may result in hypotension, hypercarbia, hypoxemia and loss of the airway.
1337 1338		<ul> <li>Movement may cause disconnection from ventilator support.</li> <li>Movement may result in accidental exturbation</li> </ul>
1339		<ul> <li>Movement may result in accidental extubation.</li> <li>Tachycardia and other dysrhythmias have been associated with transport.</li> </ul>
1340		Equipment failure can result in inaccurate data or loss of monitoring capabilities.
1341		Loss of oxygen supply may lead to hypoxemia.
1342		
1343	Contra	indications
1344		
1345 1346		Inability to provide adequate oxygenation and ventilation during transport either by manual ventilation or mechanical ventilation.
1347		Inability to maintain acceptable hemodynamic performance during transport.
1348		Inability to adequately monitor the patient's cardiopulmonary status during transport.
1349		Inability to maintain airway control during transport.
1350 1351		Transport should not be undertaken unless all the necessary members of the transport team are present.
1352 1353	Respira	ntory Care Resources
1354	_	
1355 1356		ial resuscitator with in-line pressure manometer, PEEP valve, and mask must accompany the during transport.
1357	iNOMAX DSir	
1358	Medfus	sion 3500
1359	Aeroge	n Aeroneb
1360	Portable oxygen supply ['e' cylinder(s)] of adequate volume.	
1361	Viasys	Vela Transport ventilator and patient circuit.
1362		
1363		
1364	Proced	lure
1365		
1366 1367		The RCP will gather all required respiratory care resources and bring the equipment to the bedside.

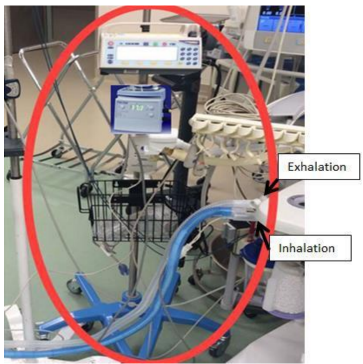
1368		The RCP will set-up and test the transport ventilator according to department procedure.
1369		* May use Vela Pole mount system or Stand Alone pole on wheels for Aerogen/Medfusion system.
1370		T <mark>he RCP will place in line the Pulmonary Vasodilator system in use per departmen</mark> t guidelines.
1371		<ul> <li>iNO Transport</li> </ul>
1372		<ul> <li>Ensure adequate supply of gas (&gt;1000 psi)</li> </ul>
1373		<ul> <li>No HME in line</li> </ul>
1374		<ul> <li>Injector Module and sampling line inserted per departmental guidelines</li> </ul>
1375		o iEpo
1376		<ul> <li>Ensure adequate supply of Epoprostenol (minimum 30 cc's)</li> </ul>
1377		<ul> <li>No HME in line</li> </ul>
1378		<ul> <li>Nebulizer placed post filter on inspiratory limb</li> </ul>
1379		<ul> <li>Expiratory filter in place</li> </ul>
1380		<ul> <li>* May use Vela Pole mount system or Stand Alone system on wheels.</li> </ul>
1381		The RCP will adjust the transport ventilator to provide an adequate level of ventilatory support
1382		• The RCP should duplicate the patient's existing ventilator settings whenever possible.
1383		<ul> <li>The RCP will insure that an appropriate respiratory rate is set (minute ventilation)</li> </ul>
1384		whenever sedation may be administered during the transport procedure to avoid
1385		hypoventilation, hypercapnia, and hypoxemia.
1386 1387		The RCP will ensure the airway is secure and in proper position before leaving the unit.
1388		• The RCP will monitor the patient during transport to insure artificial airway stability and
	_	patency.
1389 1390		Prior to leaving the unit the RCP will assess the transport ventilator's ability to provide an
		adequate level of ventilation support by observing the monitored
1391		<ul> <li>Exhaled tidal volume</li> <li>Minute volume</li> </ul>
1392		<ul> <li>Minute volume</li> </ul>
1393		<ul> <li>Respiratory rate.</li> </ul>
1394 1395		<ul> <li>SpO2</li> <li>ETCO2</li> </ul>
	_	
1396 1397		On arrival to destination, the RCP will review all above monitored parameters and verify stability of values and system.
	_	
1398		If the patient is scheduled to undergo a procedure in the operating suite, the RCP may return to
1399		his/her assigned area after verbal hand off with the Attending Anesthesiologist.
1400		• Hand off will include a review of the pulmonary vasodilator system in use, the
1401		mechanical ventilator, and the procedure for ordering inhaled epoprostenol from OR
1402		pharmacy.
1403 1404		• The RCP will provide the Anesthesiology team with his/her pager/phone number so that
		he/she may be contacted.
1405		Desumentation of two per extractly be done in the EMAD per dementer out at a done level
1406		Documentation of transport will be done in the EMAR per department standards.
1407		
1408		
1409		
1410		
1411		
1412		
1413		
1414		
1415		
1416		
1417		
1418		page 34

1419 1420		REFERENCES
1421 1422	1.	AARC Clinical Practice Guideline: In-Hospital Transport of the Mechanically Ventilated Patient – 2002 Revision and Update. Resp Care 2002; 47(7):721-723

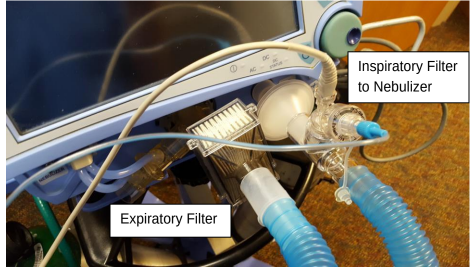
# **OR Support for Epoprostenol**

1423

1424 1425	The use of inhaled Epoprostenol is moving forward into the OR. We have supported 5 cases thus far and an additional 2 cases that were on inhaled epoprostenol and required to return to the OR.			
1425	To continue this support, please note the following and contact a supervisor or J Cappiello when			
1427	inhaled epoprostenol in the OR is requested.			
1428				
1429		New OR Case requiring Epoprostenol		
1430	1.	Contact Supervisor/J Cappiello to obtain F&P heater		
1431	2.	Obtain Medfusion/Aerogen delivery unit, 3 Aerogen nebulizers, circuit tee and infusion		
1432		tubing, heated wire circuit, and 1 L sterile water for humidification.		
1433	3.	Proceed to OR requesting unit.		
1434	4.	Place Unit between OR bed and anesthesia machine and set up circuit from anesthesia		
1435		inspiratory port, to humidifier, to patient and return to exhalation port, (as in the PB840).		
1436	5.	Inform Anesthesia to place order for Epoprostenol Inhalation		
1437	6.	If Epoprostenol available, set up syringe pump assembly. If Epoprostenol not available,		
1438		instruct anesthesia to call when Epo is available.		
1439	7.	Review circuit, delivery system, how to monitor neb, troubleshooting and dosing (IBW)		
1440		with Anesthesia.		
1441	8.	HME may be used until Epo and Humidity is initiated.		
1442	9.	Provide Anesthesia with RCS contact phone number and request to be called for Epo		
1443		initiation		
1444	10.	. Transport of this patient from the OR will occur with a transport ventilator		
1445				



1446		
1447		<b>Return to OR from CTICU on Epoprostenol</b>
1448		
1449	1.	Contact Supervisor/J Cappiello
1450	2.	Consult with Anesthesia for duration of case and intra-operative ventilation plans
1451	3.	Place patient on transport ventilator with inhaled epo as per policy. No HME
1452	4.	Place two additional nebulizers in delivery unit basket
1453	5.	Ensure Epoprostenol volume is 40 cc or greater
1454	6.	Transport patient to OR on transport ventilator
1455	7.	Review delivery system, how to monitor, trouble shooting and dosing (IBW) with
1456		Anesthesia team
1457	8.	If transport ventilator will be used for the case, review ventilator with Anesthesia team
1458		No HME.
1459	9.	Provide Anesthesia with RCS contact phone number
1460	10.	Transport of this patient from the OR will occur with a transport ventilator
1461		
1462		



- 1464 Inhaled Nitric Oxide Protocol for the Adult Lung Transplant Patient in the Cardiothoracic Intensive Care1465 Unit
- 1466
  1467 This protocol sets forth the standards for the use of inhaled nitric oxide (iNO) for the Lung Transplant
  1468 Patient in the Adult Cardiothoracic Intensive Care Unit (CTICU) by Respiratory Care Practitioners.

### 1469 1470 **PATIENT POPULATION:**

1471

1481

1484

1486

1472 Lung Transplant patients receiving inhaled Nitric Oxide (iNO) for pulmonary vasodilator therapy.

#### 1473 1474 **DESCRIPTION**:

1475
1476 INO (nitric oxide gas) is an odorless, colorless gas administered by inhalation. Nitric oxide, the active
1477 substance in INO, is a vasodilator and when inhaled, vasodilation is limited to the pulmonary
1478 vasculature.

## 1479 1480 **INDICATIONS:**

Lung transplant patients located on 7 West for the management of pulmonary hypertension, right
 ventricular dysfunction, or refractory hypoxemia.

## 1485 **ASSESSMENT OF RESPONSE**:

1487 The desired response to iNO is a decrease in pulmonary artery pressure, improved hemodynamics, 1488 and/or improved arterial oxygenation.

1489			
1490	PRECAUTIONS AND SIDE EFFECTS:		
1491 1492 1493	Hypoxemia secondary to Methemoglobinemia		
1494 1495	Airway inflammation due to elevated NO <sub>2</sub> >1.5PPM. NO <sub>2</sub> is a nitric oxide byproduct - not to be mistaken with nitrous oxide (N <sub>2</sub> O), an anesthetic gas).		
1496 1497 1498	EQUIPMENT:		
1499	INOmax DS IR		
1500 1501 1502	PROCEDURE and DOSING STRATEGIES		
1503 1504	The RCP will follow established departmental standards per INOmax DS IR policy for set up and administration.		
1505	□ INO therapy will be administered on provider order.		
1506	$\Box$ The dose range is 0.5-20 PPM with the usual starting dose of 20 PPM.		
1507 1508 1509 1510 1511 1512 1513 1514 1515 1516 1517 1518 1519 1520 1521	<ul> <li>INO therapy will be administered by the INOmax® DS IR system with the following equipment:         <ul> <li>Viasys Avea</li> <li>Puritan Bennett 840</li> <li>Viasys Vela</li> <li>SensorMedics 3100B</li> <li>Optiflo Humidification System</li> <li>Nasal cannula</li> </ul> </li> <li>Incremental dose adjustments should be done by doubling current dose (not to exceed 20 PPM). Decrements should be done by halving the current dose.</li> <li>The following parameters will be monitored to assess dose response: PA<sub>sys</sub>, PA<sub>dias</sub>, PA<sub>mean</sub>, CVP, CI, POx, SVO<sub>2</sub></li> <li>iNO can be administered during invasive or noninvasive ventilation, Nasal Cannula, and High Flow Humidity delivery.</li> </ul>		
1522	Performed twice a shift and at each dosing change to include:		
1523	<ul> <li>NO Dose set</li> </ul>		
1524	<ul> <li>NODosemonitored</li> </ul>		
1525	<ul> <li>NO2monitored</li> </ul>		
1526	<ul> <li>Right Heart hemodynamic parameters, SVO2, Pulse Oximetry</li> </ul>		
1527	<ul> <li>Tankpressure</li> </ul>		
1528	Weaning will follow below guidelines.		

1529 1530 WEANING 1531 1532 GENERAL COMMENTS: Right Heart hemodynamics include: SVO2, CI, PA sys/dia, CVP. There is a 1533 potential for rebound increase in pulmonary vasoconstriction and hypoxemia when iNO is abruptly 1534 discontinued. Weaning of iNO will commence following communication and consent of the Provider 1535 team. Prior to each dosing change, Right Heart hemodynamics will be assessed and documented in the 1536 comment section of the Nitric Flowsheet. In general, the Right Heart parameters should meet the 1537 following criteria before weaning or discontinuation: SvO2 > 60, CVP < 15, CI > 2.0, and adequate 1538 oxygenation. Pulmonary arterial pressure values will vary but should remain stable during weaning and 1539 trial off. 1540 1541 1. Obtain consent for weaning commencement from provider team. 1542 Initiate weaning if non-ECMO, stable 2 hours post-op and no bleeding (chest tube output <</li> 1543 100 cc/hr) and hemodynamically stable. Reassess every 2 hours. 1544 3. After initiation of weaning, one or more of the following conditions warrant a return to the 1545 last dose and notification of the provider team. Provider may choose to resume wean in 1 hour or halt weaning at current dose until further discussion. 1546 1547 □ PF ratio 200 or less 1548 □ PaO2 <70mmHg 1549 □ FiO2>0.40 1550 □ pulmonary hypertension (MPAP>30 mmHg) 1551 reduced SvO2 < 60 % 1552 increased CVP by more than 5 mm Hg or CVP value is > 15 mm Hg 1553 reducedCI<2.0duringanydosingdecrease 1554 1555 1556 1557

1558

Set Dose	Action
20	Dosage rate at start of wean
10	Observe Right Heart hemodynamics/oxygenation and decrease dose to 10 ppm only and allow stabilizing for 60 min.
5	Observe Right Heart hemodynamics/oxygenation and decrease dose to 5 ppm only and allow stabilizing for 60 min.
1	Observe Right Heart hemodynamics/oxygenation and decrease dose to 1 ppm only and allow stabilizing for 60 min.
0.5	Observe Right Heart hemodynamics/oxygenation and decrease dose to 0.5 ppm only and allow stabilizing for 60 min.
Off	If Right Heart hemodynamics/oxygenation are stable, notify Provider for Discontinue of iNO and observe for 60 min before taking down setup

1559		
1560	DISCO	ONTINUATION:
1561		
1562	1.	Once weaning protocol has been completed, notify the provider team of hemodynamic stability
1563		with iNO off after 1 hour.
1564		
1565	2.	If rebound worsening of pulmonary hypertension, reduced SvO2 < 60, increased CVP by more
1566		than 5mmHg or CVP value is > 15 mmHg, reduced Cl < 2.0, or hypoxemia occurs during any
1567		dosing decrease, return to last administered dose, and notify provider team. Provider may
1568		choose to resume wean in 1 hour or halt weaning at current dose until further discussion.
1569		
1570		<u>REFERENCES</u>

- 1571
- 1572 Inhaled Epoprostenol Protocol for the Adult Lung Transplant Patient in the Cardiothoracic Intensive Care
   1573 Unit
   1574
- 1575
  1576 This protocol sets forth the standards for the use of inhaled epoprostenol (iEPO) for the Lung Transplant
  1577 Patient in the Adult Cardiothoracic Intensive Care Unit (CTICU) by Respiratory Care Practitioners.

### 1578 1579 **PATIENT POPULATION:**

1580

1591

1596

1581 Lung Transplant patients receiving inhaled epoprostenol (iEPO) for pulmonary vasodilator therapy. 1582

# 1583 **DESCRIPTION**:

1584
1585 Epoprostenol (Veletri) is a naturally occurring prostaglandin that serves as a potent vasodilator and is an
effective inhibitor of platelet aggregation. Aerosolized epoprostenol is used as a selective pulmonary
vasodilator when administered by inhalation, it has been shown to improve oxygenation, reduce
pulmonary shunt, lower pulmonary artery pressure and pulmonary vascular resistance.

### 1589 1590 **INDICATIONS**:

Lung transplant patients located on 7 West for the management of pulmonary hypertension, rightventricular dysfunction, or refractory hypoxemia.

### 1594 1595 ASSESSMENT OF RESPONSE:

1597 The desired response to iEPO is a decrease in pulmonary artery pressure, improved hemodynamics, and/ 1598 or improved arterial oxygenation.

1599		
1600	PRECA	UTIONS AND SIDE EFFECTS:
1601		
1602		IEPO has fewer adverse effects than intravenous administration.
1603		Systemic effects such as hypotension, nausea, flushing, headache, and dizziness are rare except
1604		at very high doses. If hypotension should occur when initiated, immediately change to a lower
1605		dose.
1606		IEPO has an alkaline pH, it can be an irritant when inhaled, causing severe coughing.
1607		Abrupt withdrawal of iEPO can cause rebound pulmonary vasoconstriction and hypoxemia, but
1608		this is rare.
1609		
1610	EQUIP	MENT:
1611	_	
1612		Medfusion 3500 Infusion pump
1613		Aeroneb Pro-xcontrol unit, Aeroneb Pro-x nebulizer with Tadapter, Aerogen Tubing Set
1614		50mLEpoprostenolsyringepreparedbypharmacy
1615		Hydroscopic filters to be placed between expiratory limb and ventilator exhalation filter with
1616		Q4H and PRN change (for invasive ventilation)
1617		
1618	PROCE	DURE and DOSING STRATEGIES
1619		The DOD will follow entrolder and an entroleter device new Energy setup of a start of an entrol of the setup and
1620 1621		The RCP will follow established departmental standards per Epoprostenol policy for set up and administration.
1622		The usual dosage range is 5-50 ng/kg/min based on ideal body weight. Do not use a dose greater
1623		than 50ng/kg/min.
1624		The usual starting dose of iEPO is 50 ng/kg/min.
1625		Dosage adjustments should be made in increments/decrements according to the weaning
1626		protocol outlined below.
1627		The initial dose of 50ng/kg/min is at the upper end of the dose range used in most clinical
1628		studies.
1629		Assess the response to therapy within 30-60 minutes of initiation.
1630		After 4 hours of clinical stability, consideration should be given to dose reduction.
1631		At high doses, there is a potential for systemic effects, which results in systemic hypotension. If
1632		this is suspected, the dose should be lowered.
1633		There is a potential for rebound pulmonary vasoconstriction and hypoxemia when iEPO is
1634		abruptly discontinued, but it is probably less than that for inhaled nitric oxide.
1635		Be prepared to increase the FIO2 and support hemodynamics during discontinuation. If rebound
1636		occurs, it may be helpful to wean the dose slowly before discontinuation.
1637		IEPO therapy can be administered through the following equipment:
1638		<ul> <li>Viasys Avea</li> </ul>
1639		<ul> <li>Puritan Bennett 840</li> </ul>
1640		<ul> <li>Viasys Vela</li> </ul>
1641		o V60
1642		<ul> <li>Optiflo Humidification System</li> </ul>
1643		$\circ$ Nasal cannula

1644	
1645	<ul> <li>Venturi Mask</li> </ul>
1646	The following parameters will be monitored to assess dose response: PA <sub>sys</sub> , PA <sub>dias</sub> , PA <sub>mean</sub> , CVP,
1647	CI, POx, SVO <sub>2</sub>
1648	DOCUMENTATION
1649 1650	<ul> <li>A Respiratory Care Assessment and type of medication administration will be documented twice a shift</li> </ul>
1651	$\Box$ eMAR documentation will occur at each syringe change and at each dosing change.
1652 1653	<ul> <li>Right Heart hemodynamic parameters, SVO2, Pulse Oximetry will be entered into the comment section</li> </ul>
1654	
1655 1656	WEANING
1657	GENERAL COMMENTS: Right Heart hemodynamics include: SVO2,CI, PA sys/dia, CVP. There is a
1658	potential for rebound increase in pulmonary vasoconstriction and hypoxemia when iEPO is abruptly
1659	discontinued. Weaning of iEPO will commence following communication and consent of the Provider
1660	team. Prior to each dosing change, Right Heart hemodynamics will be assessed and documented in the
1661	comment section of the eMAR. In general, the Right Heart parameters should meet the following criteria
1662	before we an ing or discontinuation: SvO2 > 60, CVP < 15, CI > 2.0, and a dequate oxygenation. Pulmonary the standard equate oxygenation and the standard equation of the standard equation of the standard equation and the standard equation and the standard equation of the standard equation of the standard equation of the standard equation and the standard equation of
1663	arterial pressure values will vary but should remain stable during weaning and trial off.
1664	
1665	1. Obtain consent for weaning commencement from provider team.
1666	2. Initiate weaning if non-ECMO, stable 2 hours post-op and no bleeding (chest tube output <
1667	100 cc/hr) and hemodynamically stable. Reassess every 2 hours.
1668 1669	3. After initiation of weaning, one or more of the following conditions warrant a return to the
1670	<u>last dose</u> and notification of the provider team. Provider may choose to resume wean in 1 hour or halt weaning at current dose until further discussion.
1670	
1671	
1672	□ PaO2 <70mmHg □ FiO2>0.40
1673	$\square$ pulmonary hypertension (MPAP>30 mmHg)
1675	$\square$ reduced SvO2 < 60 %
1676	increased CVP by more than 5 mm Hg or CVP value is $> 15$ mm Hg
1677	<ul> <li>reduced CI&lt;2.0 during any dosing decrease</li> </ul>

Set Dose	Action
50	Dosage rate at start of wean
45	Observe Right Heart hemodynamics and decrease dose to 45ng/kg/min only and allow stabilizing for 1 hour.
35	Observe Right Heart hemodynamics and decrease dose to 35ng/kg/min only and allow stabilizing for 1 hour.
25	Observe Right Heart hemodynamics and decrease dose to 25ng/kg/min only and allow stabilizing for 1 hour.
15	Observe Right Heart hemodynamics and decrease dose to 15ng/kg/min only and allow stabilizing for 1 hour.
Off	If Right Heart hemodynamics are stable, notify Provider for Discontinue of Epoprostenol and observe for 1 hour before taking down setup

# **DISCONTINUATION:** 1684

1. Once weaning protocol has been completed, notify the provider team of hemodynamic stability with iEPO off after 1 hour.

## If rebound worsening of pulmonary hypertension, reduced SvO2<60, increased CVP by more than 5mmHg or CVP value is > 15 mmHg, reduced CI < 2.0, or hypoxemia occurs during any dosing decrease, return to last administered dose, and notify provider team. Provider may choose to resume wean in 1 hour or halt weaning at current dose until further discussion.

## **REFERENCES**

1694	INSPIRE-FLO Research Summary (copied from IRIS) Version 1.4
1695	Primary study objectives
1696 1697 1698	Evaluate iNO and iEPO in order to determine if they have similar impact on clinical outcomes in end- stage lung disease patients undergoing lung transplantation and end-stage heart failure patients undergoing LVAD implantation or heart transplantation.
1699	Perform a cost analysis to evaluate the average per-patient cost to use each drug.
1700	Sub-Study
1701 1702 1703 1704	The purpose of this sub-study is to understand the role of RV muscle bioenergetics using targeted metabolite profiling, which include acylcarnitines, amino acids, and ceramides, in order to (i) identify the pathway through which iPVD therapy improves outcomes and affects RV mitochondrial fatty acid utilization, and (ii) identify plasma metabolites that predict responsiveness to iPVD therapy.
1705	Secondary study objectives
1706	Sub-Study
1707 1708 1709 1710	1.Circulating metabolic biomarkers will identify heterogeneity of response to iPVD therapy, by reporting on this underlying RV muscle's mitochondrial fatty acid utilization for energy production.
1711	Standard Research Summary
1712	Purpose of the Study
1713	Objectives & hypotheses to be tested
1714 1715 1716 1717 1718 1719 1720 1721	1. Aim I – Clinical Trial Investigation. In order to utilize Inhaled Epoprostrenol (iEPO, Veletri®, Actelion Pharmaceuticals, South San Francisco, CA, USA) as an acceptable alternative to Nitric Oxide (iNO, INOMAX®, Mallinkrodt Pharmaceuticals, St. Louis, MO, USA) in adult patients, we propose a randomized, prospective, double-blinded trial in the cardiothoracic surgical population, which will evaluate the primary hypothesis that these two medications will have similar efficacy in pulmonary vasodilation and a similar impact on clinical outcomes in end- stage lung disease patients undergoing lung transplantation and end-stage heart failure patients under durable LVAD implantation or heart transplantation (Table 1).
1722 1723 1724 1725 1726	2. Aim II – Cost-Capture Analysis. There will be a parallel prospective cost-capture analysis designed to precisely acquire the expenses that each drug incurs per patient averaged across all patients randomized to that drug.
1727	Sub-Study
1728 1729 1730 1731 1732 1733	Central hypothesis - Patients who are refractory to changes in pulmonary vascular tone after iPVD therapy will also display reduced right ventricular (RV) mitochondrial fatty acid utilization through increased lipid infiltration of right heart muscle.
1734	Page 46

1739 1740

 Secondary Hypothesis - Circulating metabolic biomarkers will identify heterogeneity of response to iPVD therapy, by reporting on this underlying RV mitochondrial fatty acid utilization.

## 1741 Background & Significance

1742 Introduction. Inhaled Nitric Oxide (iNO) is a selective pulmonary vasodilator (PVD) with FDA-approval in 1743 the neonatal population alone. In adult patients, iNO is used off-label to treat pulmonary hypertension, 1744 right ventricular (RV) failure, and ventilation-to-perfusion mismatch. Adult patients who undergo 1745 durable LVAD implantation (e.g. Heartware®, Heartmate 2®, or Heartmate 3®) or cardiac transplantation 1746 for end-stage heart failure or those that have endured lung transplantation as a result of end-stage lung 1747 disease, compose the largest subpopulation which receives PVD therapy at Duke University Hospital. 1748 Intravenous Epoprostenolis FDA approved for adult patients with pulmonary hypertension and is the 1749 only agent which has displayed mortality benefit in these patients. The inhaled formulation of 1750 Epoprostenol (iEPO) was developed in order to maintain efficacy and avoid the systemic side effects of 1751 vasodilation and thrombocytopenia. Inhaled iEPO is used off-label in our cardiothoracic surgical patients 1752 for new-onset perioperative pulmonary arterial hypertension (PAH), known preoperative PAH, RV 1753 dysfunction with LVEF > 35-40%, and promotion of ventilation to perfusion matching through alveolar 1754 deposition of the prostanoid compound and vasodilation of the intimately associated intra-acinar 1755 pulmonary arteries. This vasodilation can decrease pulmonary vascular resistance and can improve 1756 oxygenation while avoiding systemic effects commonly seen in the intravenous formulation. iEPO has 1757 been introduced in the cardiothoracic operating rooms (OR) and ICU as a cost-conscious alternative 1758 medication to iNO. iEPO may display an equivalent efficacy profile to iNO for pulmonary vasodilation 1759 and oxygenation and have a similar impact on clinical outcomes. For the purposes of this writing, 1760 thoracic transplantation will refer to both heart and lung transplantation.

1761 Pharmacology. There are 3 major pathways that affect pulmonary vascular tone: 1) Nitric oxide 1762 (vasodilatory), 2) Prostaglandin (vasodilatory), and 3) Endothelin (vasoconstrictive) pathways. During 1763 cardiothoracic operations, particularly transplantation and LVAD surgery, there is an appreciable 1764 imbalance in these pathways, which favors vasoconstriction. iNO administration, exerts its mechanism 1765 of pulmonary vasodilation and ventilation-to-perfusion matching through exogenous NO delivery and 1766 iEPO applies a similar mechanism via exogenous prostacyclin delivery. Both agents are delivered through 1767 mechanical ventilation to ventilated alveoli in order to promote gas exchange at the capillary bed. Both 1768 inhaled medications are desirable in this population due to pulmonary selectivity, absence of systemic 1769 vasodilation, as well as fast onset (5-10 seconds for iNO and 30-60 seconds for iEPO) and quick titration 1770 owing to short-half lives (10-20 seconds for iNO and 1-2 minutes for iEPO). There is no decision tree 1771 involved in the use of iNO vs iEPO except for that patient's known drug allergies which may preclude use 1772 of one inhaled agent in favor of the other. Of note, endothelin antagonists (e.g. bosentan), which are 1773 not part of our perioperative standard practice, are PO medications which require reliable 1774 gastrointestinal absorption that may not be present during high-dose inotropic support, and are not 1775 readily titrated to effect as are the inhaled PVD, iNO and iEPO.

1776 Contraindications and Adverse Effects. Absolute indications for iNO in favor of iEPO are due to
 1777 prostaglandin allergy leading to anaphylaxis (extremely rare) or if the patient is pregnant due to risk for
 1778 labor induction as a result of prostacyclin agonism. Routine pregnancy testing is performed in the

INSPIRE-FLO Research Summary (copied from IRIS) Version 1.4

1779

preoperative setting in line with established preoperative anesthesia testing criteria. Parturients rarely
present for thoracic transplantation or LVAD implantation. There are no absolute contraindications to
iNO therapy in adult patients but the iNO delivery device system routinely measures the toxic
metabolite of iNO, nitrogen dioxide (NO2), which can lead to hypoxemia during metabolite
accumulation. Additionally, methemoglobinemia (MetHb) is another rare adverse occurrence of
prolonged iNO administration and MetHb levels are measured during arterial blood gas analysis.

1786 Preliminary retrospective study supporting noninferiority hypothesis. In a retrospective study of 51 adult 1787 cardiothoracic surgical patients (all-comers, including thoracic transplantation, durable LVAD 1788 implantation, and non-transplant and non-LVAD cardiac surgical patients), requiring pulmonary 1789 vasodilation, our group illustrated similar efficacy between the use of iEPO and iNO with respect to 1790 optimizing RV hemodynamic variables, including pulmonary vasodilation and mixed venous oxygenation 1791 (Table 2). During this investigation, iNO was initiated in the operating room (OR) and continued during 1792 transport and into the ICU. While in the ICU, postoperative hemodynamic stability was achieved within 2 1793 hours and iNO was transitioned to iEPO over 30 minutes in order to provide continuous inhaled 1794 pulmonary vasodilation and allow the patient to self-control during medication cross-over between iNO 1795 and iEPO. Clinical variables were followed at 5-minute intervals for 1 hour after transition to iEPO. No 1796 statistically significant differences were seen in hemodynamic variables during this transition (Table 2). 1797 The small sample size and retrospective design, however, incorporated several confounding variables 1798 that could not be controlled and prospective data was deemed necessary to achieve reliable conclusions 1799 by evaluating clinical outcomes in order to change clinician practice patterns. Other investigations have 1800 demonstrated equivalence in hemodynamic variables, mixed venous oxygenation, and ventilation-to-1801 perfusion matching when delivery of iNO was compared with iEPO. These studies were, however, also 1802 retrospective or inadequately powered to rely on conclusions related to outcome measures.

1803 The large cost differential between these two agents remains an important concern for the health 1804 system: iNO is approximately 8-fold more expensive than iEPO, according to preliminary estimates based 1805 on PVD usage. Previous reports have estimated the cost of iNO administration to be between \$95.00 -1806 \$115.00 perhour during medication delivery. The cost, however, has not precisely captured the time 1807 required to assemble the iNO delivery system as well as resources utilized to breakdown this setup into 1808 individual components following termination of delivery. The cost of iEPO delivery is captured at \$14.83 1809 per hour, which includes solution compounding by pharmacy as well as processing for delivery and 1810 nebulization by respiratory care services. Additionally, the iEPO delivery-system setup is a one-time, 1811 fixed cost for the duration of administration. Similar secondary resource utilization capture for iEPO is 1812 required for accurate cost comparison between these two agents.

- 1813 1814
- 1815 Sub Study

Acute right-sided heart failure (aRHF) strongly predicts the incidence of early death after left ventricular
 assist device placement (Soliman Oll) or heart transplantation (Taghavi S) for the surgical treatment of
 advanced left-sided heart failure. Although inhaled pulmonary vasodilator (iPVD) therapy is the
 mainstay for vascular afterload reduction in aRHF, more than 40% of patients may be refractory to
 treatment and display persistently elevated pulmonary vascular tone without improvement in right
 ventricular (RV) muscle contractility. Healthy RV muscle contraction utilizes long-chain fatty acids,
 delivered to mitochondria by acylcarnitine molecules, for efficient energy production. Prior work in

1824 patients with aRHF and pulmonary hypertension has identified derangements in fatty acid transport into 1825 mitochondria by acylcarnitines. Notably, lipid infiltration in RV muscle occurs during aRHF with 1826 overexpression of ceramide metabolites (biomarkers of lipotoxicity). (Brittain EL) In circulating plasma, 1827 long-chain fatty acid acylcarnitine (LC FA AC) metabolites are displaced from RV muscle cells and serve 1828 as biomarkers for aRHF. (Brittain and Luo) Reduced levels of LC FA AC are illustrated in such patients who 1829 respond to vasodilators by lowering pulmonary vascular tone and this response is associated with 1830 improved survival. (Rhodes CJ) Members of my mentorship committee have shown that plasma LC FAAC 1831 levels decrease to pre-left HF values after left ventricular assist device placement. (Ahmad T) However, 1832 the molecular pathways underlying aRHF after surgery remain unknown and the clinical utility of 1833 complimentary plasma biomarkers in guiding therapy has not been defined. Therefore, there is a critical 1834 need to identify the underlying metabolic aberrancies in RV muscle of cardiac surgical patients with 1835 postoperative aRHF refractory to iPVD therapy. Without such information, novel personalized 1836 therapeutic targets for aRHF after surgery will remain limited

## 1837 Design & Procedures

1838 Aim I – Development of a Definitive Clinical Trial Investigation.

1839 1. Randomization and Double-Blinding. The clinical research unit (CRU) will receive preoperative 1840 notification of lung and heart transplantation patients by reviewing the transplant waitlist. Preoperative 1841 notification of LVAD implantation will be done by the review of the cardiothoracic surgical schedule. 1842 Using a 50% randomization process utilized and established by the CRU at Duke University Hospital, 1843 each eligible patient will be randomized to receive either iNO or iEPO. The primary endpoint data will be 1844 collected and documented in an electronic data capture system during the period of time the patient, 1845 clinical care team, and study team are blinded. Primary endpoint data collection will be complete prior 1846 to the subjects' discharge from the ICU, at which point the unblinding will occur. Since primary endpoint 1847 data collection will occur during the blinded period, the potential for bias will be substantially 1848 minimized.

1849 2. Measured Outcomes. The primary endpoint for the comparison of efficacy in the Lung Transplant 1850 population will be the incidence of Grade 3 Primary Graft Dysfunction (PGD). This is defined by the 1851 International Society of Heart and Lung Transplantation (ISHLT) as severe hypoxemia with a PaO2-to-1852 FiO2 ratio < 200 or the presence of venovenous extracorporeal membrane oxygenation (VV ECMO) at an 1853 time-point within the first 72 hours after lung transplantation. The primary endpoint for the comparison 1854 of efficacy in LVAD patients will be incidence of moderate or severe RV failure according to Interagency 1855 Registry for Mechanically Assisted Circulatory Support (INTERMACS) scoring. The primary endpoint for 1856 the comparison of efficacy in the heart transplant subset will be the incidence rate of RVAD insertion. 1857 Secondary endpoints related to clinical outcomes for all populations will be duration of postoperative 1858 mechanical ventilation, , ICU Length of Stay (LOS), hospital LOS, incidence of acute kidney injury, 1859 incidence of in-hospital mortality, as well as postoperative mortality at 30-days, 90-days, and 1-year 1860 after operation (Table 1).

1861 Aim II – Cost-Capture Analysis.

In parallel with the design & procedures of Aim I, the cost capture analysis component will be essential
 in order to better gauge the cost due to duration of administration (variable cost) according to each
 inhaled PVD. Established clinical criteria specific to each group (lung transplantation vs. heart

transplantation/LVAD implantation) have been developed to determine the inception of protocolized
 PVD weaning. Weaning medications according to established protocols will allow for accurate
 interpretation of the comparative length of therapy between iNO and iEPO and help prevent erroneous
 PVD usage without criteria for discontinuation. Secondary resource utilization will be documented by
 respiratory care services and itemized cost sheets will be developed.

1871 Sub-Study

1865

1872 This sub study will leverage the target enrollment of 224 patients undergoing left ventricular assist 1873 device placement or heart transplantation in the parent study. Furthermore, cardiac tissue is collected in 1874 collaboration with the Duke Human Heart Repository (IRB PRO#00005621) and data will be used to test 1875 the causal relationship between RV myocardial fatty infiltration and plasma elevation of LC FA 1876 acylcarnitines in patients refractory to iPVD who develop acute RHF. This tissue and plasma is already 1877 being collected. In patients receiving a heart transplant, one RV tissue core sample will be obtained 1878 during routine, standard-of-care, post-operative endomyocardial biopsy. Core samples are the size of 1879 2mm pellets.

- 1880 Selection of Subjects
- 1881 Subject Groups

1882 Inhaled PVD therapy is administered to every patient undergoing thoracic transplantation and LVAD 1883 implantation at our institution and each patient is eligible for enrollment. Over a 3-year period (1 year 1884 for follow-up) we will prospectively enroll 200 lung transplant subjects and 224 heart transplant or LVAD 1885 implantation patients who will be informed and consented prior to their scheduled procedure. Potential 1886 subjects will be under the care of 1 or more investigators in this study. Consented subjects will be 1887 randomly assigned to 1 of 2 groups, iNO vs iEPO, to be initiated in the OR on the day of the operation 1888 based on accepted standard of practice and study protocol. Medication administration will be double-1889 blinded, such that neither the surgical nor an esthesiology teams will be notified of the inhaled agent to 1890 which the patient has been randomized. Ability to unblind the delivery system will be made available to 1891 both teams if required to preserve optimal patient care. As per our standard practice, respiratory care 1892 services will manage the initiation and maintenance of inhaled PVDs in the OR and ICU, and these 1893 personnel will be the only practitioners notified of the actual delivered medication during study blinding.

- 1894 1895
- 1896 Exclusion Criteria
- 1897 Combined Organ Transplantation (e.g., Heart-Lung, Heart-Liver, Heart-Kidney, Lung-Liver, etc.)
- 1898 · Age < 18 years old
- Pregnancy (females of child bearing potential will receive pregnancy testing prior to cardiothoracic
   surgery as a standard of care)
- 1901 · Known allergy to prostaglandin (rare)
- 1902 · Refusal of blood products due to personal or religious preference.
- 1903 . Subject is enrolled in another study protocol, which does not allow randomization of PVD therapy

INSPIRE-FLO Research Summary (copied from IRIS) Version 1.4
· Heart transplant or durable LVAD recipients with adult congenital heart disease (CHD)
Caveat: Does NOT meet exclusion criteria if the scheduled heart transplant or LVAD implantation is due to heart failure from a previous heart transplantation related to CHD, performed more than 90 days previous to the date of trial enrollment
• Patient is scheduled to undergo lung transplantation but has undergone heart transplantation in the previous 90 days
• Patient is scheduled to undergo durable LVAD implantation but has undergone heart transplantation in the previous 90 days
• Patient is scheduled to undergo heart transplantation but has undergone lung transplantation in the previous 90 days
· Patients with preoperative VV ECMO as a bridge to lung transplantation
Stopping Criteria – In the event the following criteria are met and the clinical team is in agreement, subjects will be weaned off of their iPVD per instutional standard iPVD weaning practice. If adverse events are encountered, the drug will be immediately stopped without weaning.
Venoarterial (VA) ECMO insertion remains at end of operation
VA ECMO insertion is performed postoperatively in the ICU
LVEF < 30% on echocardiogram at the end of the operation for heart and lung transplant subjects
LVEF < 30% for heart and lung transplant subjects on echocardiogram noted postoperative in the ICU
Inhaled pulmonary vasodilation is halted for reasons other than standard weaning ordered by the clinical care team
Adverse events related to the INO or EPO that affect the subject's welfare
Data Collection
Secondary measures will be hemodynamic variables (similar to those measured in Table 2) such as transesophageal echocardiographic (TEE) evaluation of RV function based on stand-of-practice protocol, intravenous administration of inotropes, serial measures of postoperative serum creatinine and GFR, resolution of elevated liver function tests (heart failure patients, illustrates improvement in RV function), incidence of thrombocytopenia (platelet count < 150 x 109/L) and trajectory of resolution, as well as ventilation-to-perfusion matching (arterial oxygen tension, PaO2; arterial carbon dioxide tension, PaCO2; and fraction of inspired oxygen, FiO2). Variables will be recorded at designated time points during the entire duration of administration – from initiation in the operating room to cessation in the ICU. These time points include: Intraoperative before surgical incision, time = 0 (initiation of PVD), 30 minutes, 2 hours, 6 hours, 12 hours, 18 hours, 24 hours, and every 6 hours up through 72 hours after initiation. These secondary measures will be obtained up through 72 hours after initiation regardless of cessation or continuation of the inhaled PVD. After 72 hours, increments of every 12 hours thereafter will be assessed if PVD administration continues. Ventilation and perfusion nuclear scans will be obtained and recorded per standard clinical practice for each group of lung transplant recipients. Established protocols with criteria for initiation of medication weaning have been created according to

INSPIRE-FLO Research Summary (copied from IRIS) Version 1.4

- 1942
- each medication based on individual pharmacokinetic properties. Once established criteria are met,
  weaning of each inhaled PVD will begin and continue until the medication is terminated according to
  standardized weaning protocols established for lung transplant patients and heart transplant/LVAD
  patients.

Subject follow up. Subject will be contacted by phone by a member of the research team and be asked a
short series of questions to assess their current medical condition and any changes since surgery at 30days (±3 days), 90-days (±5 days), and 1-year (±7 days) after surgery completion date. The phone
follow-up should take approximately 5 minutes of the subject's time. If subjects have been admitted to a
hospital outside of Duke Health after surgery they will be asked to sign an authorization of release to
provide us permission to obtain medical information related to their hospitalization.

- 1953
- 1954
- 1955 Blood Sampling

1956 Blood samples will be drawn for analysis as a part of this study. One 9 ml sample of blood will be 1957 obtained from each patient prior to the initiation of PVD therapy and stored at 4°C prior to processing. 1958 This sample will be stored for Genomic DNA analysis at the completion of this study in order to assess 1959 patients who are responders to inhaled pulmonary vasodilaton through upregulation and down 1960 regulation of notable vasoactive substances (e.g. endothelin, thromboxane, nitric oxide, prostaglandin, 1961 etc.). In addition, each subject will also be asked to sign the Genomic and Proteomic Database 1962 Repository (IRB Pro00015651) consent form, thus allowing the banking of their plasma and DNA samples 1963 as well as data to be used for future research. Participation in IRB Pro00015651 is voluntary and optional 1964 to all subjects consented in this parent study. Blood samples (7 ml each) will be drawn at 3 separate 1965 time points: 1) directly after insertion of the invasive blood pressure monitoring (arterial) line, 2) POD 1 1966 (8 to 24 hours after completion of surgery), and 3) POD7 (6 days from POD 1). In each 7ml blood 1967 sample, 3.5ml will be collected in Sodium Citrate tubes for coagulation analysis and another 3.5ml will 1968 be collected in EDTA tubes for metabolomic and proteomic analysis. Plasma will be separated from 1969 these samples and banked at -80°C for analyses of proteomic and metabolomic signatures. Up to 30ml 1970 of blood will be collected during the 12 month study participation period.

1971 Sub-Study

1972 This sub-study only pertains to LVAD and heart transplant patients who meet inclusion criteria and have 1973 consented to participate in the parent study. For heart transplant patients, the POD 7 sample collection 1974 will occur on the day of the biopsy procedure regardless of post operative day.

## 1975 Subject Recruitment and Compensation

1976Describe recruitment procedures, including who will introduce the study to potential subjects. Describe1977how you will ensure that subject selection is equitable and all relevant demographic groups have access1978to study participation (per45 CFR 46.111(a) (3)). Include information about approximately how many1979DUHS subjects will be recruited. If subjects are to be compensated, provide specific prorated amounts to1980be provided for expenses such as travel and/or lost wages, and/or for inducement to participate.

Subjects will be recruited either during the outpatient or inpatient evaluation phase, or contacted by
 phone. Recruitment may also occur on the day of the operation given the complexities of the transplant

1984 process, which may provide obstacles to earlier enrollment. After obtaining permission from the 1985 operating surgeon, surgical subjects will be screened by the study coordinator by reviewing the 1986 transplant pre-list. Prior to asking any patient for consent to participate, the patient or Legally 1987 Authorized Representative (LAR) will be approached first by the surgeon or one of the members of the 1988 surgical care team to determine if the patient or LAR is willing to consider enrollment in the study. If so, 1989 the subject or LAR will either be seen during an inpatient or outpatient visit, or be contacted by phone 1990 and informed about the study by a member of the research team. If the individual or LAR is willing to 1991 consider enrollment and does not meet exclusion criteria, then the research coordinator will present the 1992 research protocol in its entirety. During this time, the study coordinator will answer any and all 1993 questions as they arise. If the subject or LAR agrees to participate, the coordinator will ask the them to 1994 sign and date the appropriate consent form. A copy of this consent form will be given to the subject and 1995 a copy of the consent form will be added to the subject's medical record. The subject or LAR will be 1996 given the option to sign a separate consent form to allow us to store portions of the collected blood 1997 specimens and any data collected under this research study and maintain these samples and data in a 1998 database/repository (PRO00015651) for possible use in future research studies relating to surgical 1999 outcomes. In the event a LAR provides consent at the time of enrollment, the subject will be 2000 approached once they regain the ability to provide an informed consent.

Recruitment will not routinely occur on the day of the operation and most patients will be enrolled at
 least 12 hours in advance and provided at least the allowable time to review the study consent form and
 discuss their options with the PI and study personnel. There will be no direct compensation to the
 patient for recruitment.

If a subject is enrolled and randomized in this study for their LVAD implantation procedure and is later
 planned to receive a heart transplant, that previously enrolled subject is eligible to be re-enrolled. The
 following caveats apply to this subpopulation of LVAD patients:

A)Durable LVAD implantation may occur as a bridge to heart transplantation.

B) If LVAD implantation is followed by heart transplantation WITHIN 1 year following LVAD implantation,
 then data collected up through the time of heart transplantation will be recorded and valid as a patient
 in the LVAD group.

- C)Data collected on or after the date of LVAD explantation/heart transplantation for such a patient willbe considered as part of the heart transplant group.
- D) If LVAD implantation is followed by heart transplantation AFTER 1 year following LVAD implantation,
   then the 1 year follow-up period is complete and the patient may re-enter the trial as a heart transplant
   patient.
- 2017 If a subject is enrolled and randomized in this study for their durable LVAD implantation procedure and 2018 is scheduled to receive a new durable LVAD via an LVAD exchange operation, the subject is eligible to be 2019 re-enrolled.
- 2020
- 2021
- 2022 Sub-Study

2023	INSPIRE-FLO Research Summary (copied from IRIS) Version 1.4
2024 2025	Only subjects consented in the parent study will be asked to participate in the substudy and they will have opt in/out ability.
2026	Consent Process
2027	Subject's Capacity to Give Legally Effective Consent
2028 2029 2030	If subjects who do not have the capacity to give legally effective consent are included, describe how diminished capacity will be assessed. Will a periodic reassessment occur? If so, when? Will the subject be consented if the decisional capacity improves?
2031 2032 2033	Explicit (written) consent will be obtained from the patient or the patient's legal decision maker.
2034	Study Interventions
2035 2036 2037 2038 2039	Using a 50% randomization process utilized and established by the CRU, each eligible patient will be randomized to receive either iNO or iEPO, to be initiated in the OR based on accepted standard of practice at Duke University Hospital, during the clinical care of these patients.
2000	Risk/Benefit Assessment
2041 2042 2043 2044 2045 2046 2047	Include a thorough description of how risks and discomforts will be minimized (per 45 CFR 46.111(a) (1 and 2)). Consider physical, psychological, legal, economic and social risks as applicable. If vulnerable populations are to be included (such as children, pregnant women, prisoners or cognitively impaired adults), what special precautions will be used to minimize risks to these subjects? Also identify what available alternatives the person has if he/she chooses not to participate in the study. Describe the possible benefits to the subject. What is the importance of the knowledge expected to result from the research?
2048 2049 2050 2051 2052 2053 2054 2055 2056 2057 2058 2059 2060 2061	There is no direct benefit of this study to the enrolled subjects. Data gathered from this study may benefit future patients. Up to 30 ml of blood will be drawn during the 12 month study participation period. Blood sampling will be obtained, in the majority of subjects, from indwelling arterial or central venous lines inserted at the beginning of the intraoperative period as part of standard practice for these operations and there will be no additional risk to the patient for obtaining such vascular access. On rare occasion, blood sampling may be obtained from additional venipuncture sites during the postoperative period. Risks of blood sampling if obtained through venipuncture are pain, swelling, possible infection at the site of venipuncture. While these risks are minimal, the additional blood volume is highly unlikely to contribute to the patient's need for blood transfusion. To minimize any potential risk to the patient from genetic data, investigators and patients will be blinded to the individual patient's genotype. This information will not be included in the patient chart, will remain absolutely confidential, and will not be given to the patient or their family. DNA samples will be identified only by a coded number whose relation to the patient's name and other identifiers is available only to the data manager. The identity of the patient will remain anonymous in any publications which may result from this investigation.
2062	There will be no additional risks to the subjects as a result of this study. Prior to June of 2015, iNO was

 $2063 \qquad the sole option for inhaled pulmonary vaso dilation in this patient population and therefore utilized in$ 

2065 each operation for this indication. As of June 2015, iEPO was introduced for the same indications as iNO 2066 in order to serve as a cost-conscious alternative to iNO and to potentially explore a different, equally 2067 impactful pathway for clinically evident pulmonary vasodilation (as measured by Swan-Ganz catheter 2068 data and determined by transesophageal echocardiography). There are no additional risks to the patient 2069 aside from the rare adverse effects such as allergic reaction, as previously discussed. The most common 2070 side effect of iNO is hypotension. The side effects common to intravenous iEPO are nausea, vomiting, 2071 hypotension, flushing, chest pain, anxiety, dizziness, bradycardia, difficulty breathing, abdominal pain, 2072 musculoskeletal pain and tachycardia

- 2073 Costs to the Subject
- 2074 There will be no additional costs to the subjects as a result of this study

## 2075 Data Analysis & Statistical Considerations

2076 Summary statistics will be computed for demographic, clinical, and outcome variables in the form of 2077 frequencies (percentage) for categorical variables and mean (standard deviation) for continuous 2078 variables for each arm. Univariate analysis will be performed to compare the difference of each variable 2079 between treatment groups by chi-square or Fisher exact tests for categorical variables, and t-tests or 2080 Wilcoxon Rank-Sum tests for continuous variables depending on data normality. The univariate results 2081 for the outcome variables will provide information on iNO treatment effect in comparison to iEPO 2082 without taking into account other potential confounding factors. All non-outcome variables meeting p< 2083 0.15 association with treatment group will be considered for variable selection to build a multivariable 2084 regression model. For each outcome of interest, we will start with a regression model (logistic 2085 regression for binary outcomes or generalized linear model for continuous outcomes) with all variables 2086 selected from univariate analysis described above. Based on stepwise variable selection, we will 2087 determine the final set of covariates to be included in the final multivariable model to test the 2088 treatment group effect. Based on the analysis results, we will be able to understand if iNO is equivalent 2089 to iEPO (no significant difference) or significantly better or worse than iEPO (significant treatment effect) 2090 to address the efficacy of iNO for Aim 1. Several of secondary measures will be obtained over time. We 2091 will apply generalized mixed model to take into account the repeated measures over time to test for 2092 treatment effect. In the case of patients have switched to the other arm due to clinical decision, we will 2093 conduct the primary analysis based on the intent to treat (ITT) without reclassifying treatment 2094 assignment. In addition, protocol analysis, where only patients follow the protocol assignment are 2095 included will also be conducted to verify ITT results. For Aim 2 to compare cost capture analysis, the 2096 comparison of cost measures between two groups will be tested by two sample t-test.

2097 Based on recent annual operations, approximately 120 LVAD implantations, 60 heart transplantations, 2098 and 110 Lung transplantations were performed at Duke University Hospital during FY 2014 – 2015. This 2099 study has been individually powered to primary endpoints for each arm (Table 1) and the duration of 2100 study enrollment has been determined according to annual operations and sample-size calculations. We estimated sample size based on equivalence test of the incidence rates of a binary outcome (e.g. PGD 2101 2102 grade 3 (PGD-3)) of two treatment groups as an illustration. Assuming the incidence rate of PGD-3 under 2103 iEPOtreatment is 0.35 and acceptable margin of the equivalence is ±0.19, we will need 224 patients to 2104 have 80% power to detect an actual difference at 0.05 between two treatment group under this margin. 2105 This implies that the acceptable range of incidence rate for iNO treatment is from 0.21 to 0.59. Based on 2106 this estimate, we propose to enroll 200 lung transplant patients and 224 LVAD and heart transplant

- 2107
- patients (n = 424) over a period of 24 to 36 months; the exact time point for trial culmination between
   24 and 36 months will be dependent on enrollment rate. There will be a 50% randomization rate for
   each inhaled agent such that 212 patients will receive iEPO and 212 patients will receive iNO.

# 2111 Data & Safety Monitoring

- The proposal is not introducing a new medication that has not been utilized by our group and safety has
   been established for this patient population through clinical practice and medication usage. Safety will,
   however, be determined by assessing reported, rare, adverse effects of iNO (systemic hypotension,
   methemoglobinemia, and rebound pulmonary hypertension after appropriate weaning) and iEPO
   (systemic hypotension, non-surgical bleeding related to thrombocytopenia, flushing, and rebound
- pulmonary hypertension after appropriate weaning) in order to accurately monitor adverse events (AE)
   during this study. The PI will review and sign off on AE's as they occur and perform a quarterly review
   and determine if AE's are related to the study or otherwise. AE's will be reported to the IRB per HRPP
- 2120 policies.
- 2121 Stopping Rule: Subjects who meet the stopping criteria in section 4 continued to be enrolled and 2122 followed for primary outcome analysis.

# 2123 Describe Role of External Personnel:

2124 All data collected in the case report forms (CRF) will be collected by review of the subjects routine 2125 medical record documentation or during the intraoperative portion of the study. All subjects will be 2126 given a study ID in an order to maintain their identity and subject's identity will be protected and 2127 confidentially maintained. Barcodes will be affixed to each study sample collected according to the 2128 protocol. For future review, the study number and barcode will be the only identifying information 2129 associated with the subject. All paper data will be stored in a locked cabinet in the research teams office 2130 as outlined in the research data security plan. Any computerized data will be stored within the Duke 2131 University Medical Center's Database, which is password protected, and located behind Duke 2132 Computing firewalls. Only the PI and the statisticians will have access to the data obtained from these 2133 cases.