



BRONCHIOLITIS OBLITERANS SYNDROME

- Lung transplantation (LTx) recipients have 1-, 5- and 10-year unadjusted survival rates of 80%, 54%, and 32%, respectively¹.
- Most deaths after first post-transplant year are caused by Chronic Lung Allograft Dysfunction (CLAD)¹.
- Bronchiolitis obliterans syndrome (BOS) is the obstructive phenotype and most common form of $CLAD^{2}$.
- Incidence of CLAD-BOS following lung transplant is about 10% per year, with a prevalence of 50% at 5/ years³

Currently there are no proven therapies for the treatment of BOS

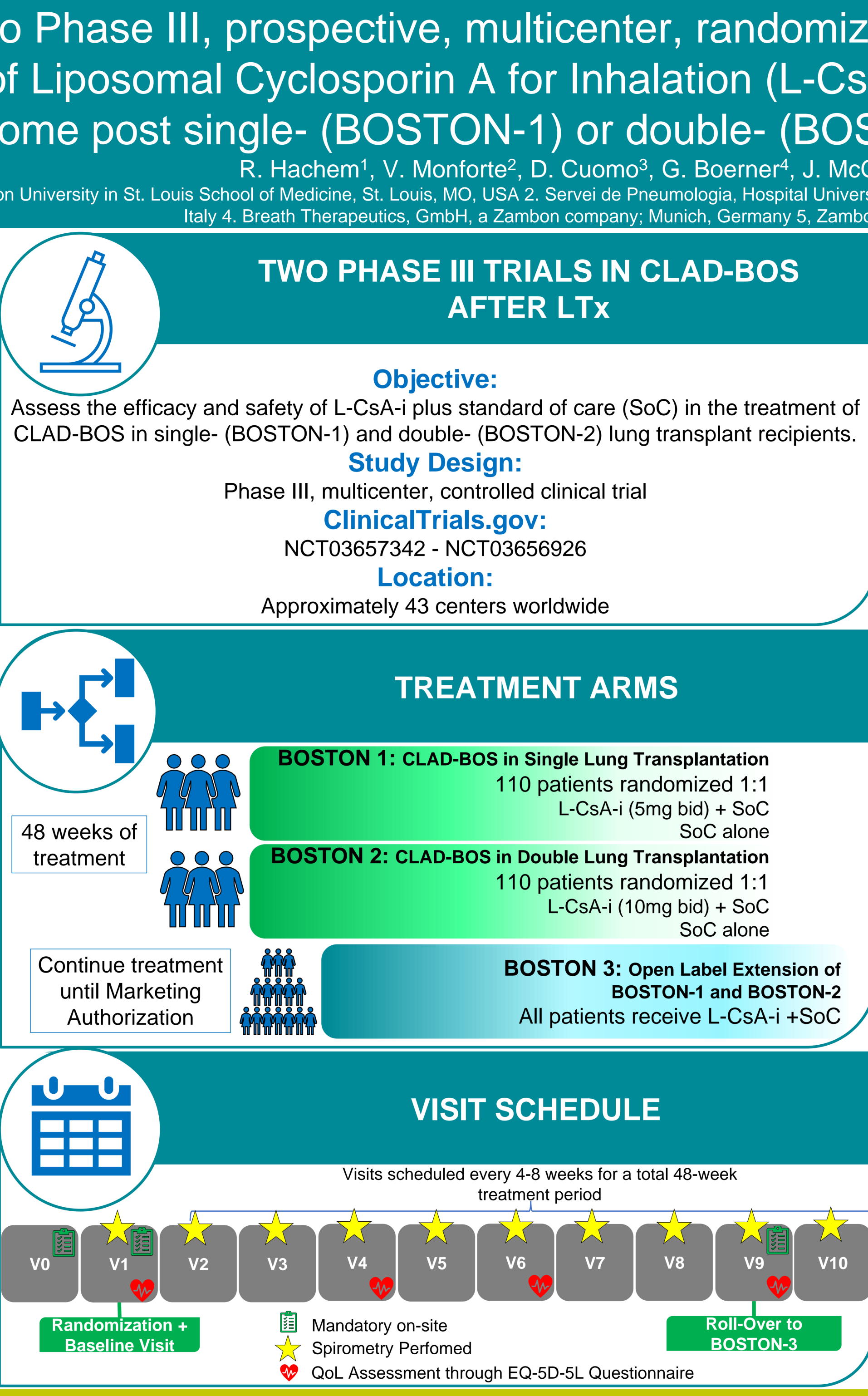


THE IDEA BEHIND THE THERAPY

- Inhaled liposomal Cyclosporine A (L-CsA-i) is an investigational formulation of cyclosporine.
- The unique formulation of L-CsA-i, administered via the investigational drug-specific eFlow® Technology nebulizer system, is designed to deliver drug directly to the small airways of the lung.
- Delivering the immunosuppressive therapy directly to the allograft, it can act locally at the site of immune activation.

REFERENCES

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Study design of two Phase III, prospective, multicenter, randomized, controlled clinical trials evaluating the efficacy and safety of Liposomal Cyclosporin A for Inhalation (L-CsA-i) in patients with Bronchiolitis Obliterans Syndrome post single- (BOSTON-1) or double- (BOSTON-2) lung transplantation R. Hachem¹, V. Monforte², D. Cuomo³, G. Boerner⁴, J. McGrain⁵, S. Prante⁴ 1. Division of Pulmonary & Critical Care, Washington Universitari Vall d'Hebron, Universitat Autònoma de Barcelona, Barcelona, Spain 3. Zambon SpA, Bresso (MI), Italy 4. Breath Therapeutics, GmbH, a Zambon company; Munich, Germany 5, Zambon USA Ltd., Cambridge, USA **KEY INCLUSION/EXCLUSION CRITERIA Exclusion Criteria: Inclusion Criteria:** Patients with BOS diagnosis defined as CLAD-BOS phenotype with: Screening FEV₁ between 85-51% of personal best FEV₁ value post-transplant **OR** phenotype) Screening FEV₁>85% of personal best FEV₁ associated with EITHER a \geq 200mL decrease in FEV₁ in the previous 12 months OR according to medical history showing BOS progression Diagnosis of CLAD-BOS must be made at least 12 months after lung transplantation and a) within 12 months prior to the screening visit **OR** b) more than 12 months from study screening and patient must have shown a decline in $FEV_1 \ge 1$ 200ml in the previous 12 months before screening, which cyclosporine A is not due to acute infection or acute organ rejection **PRIMARY ENDPOINT:** Mean change in FEV1(mL) from baseline to Week 48 **110** patients randomized 1:1 **SECONDARY ENDPOINTS:** L-CsA-i (5mg bid) + SoC Mean change in FEV₁/FVC from baseline to Week 48 SoC alone Time to Progression of BOS, defined as the earliest of the following: Absolute decrease from baseline in FEV₁≥10% or ≥200 mL and absolute decrease in FEV₁/FVC **110** patients randomized 1:1 of >5% **OR** L-CsA-i (10mg bid) + SoC Change in BOS Severity, **OR** SoC alone Re-transplantation, **OR** Death from respiratory failure **BOSTON 3:** Open Label Extension of **BOSTON-1** and **BOSTON-2** All patients receive L-CsA-i +SoC CONCLUSIONS These phase III trials will characterize the safety and efficacy of L-CsA-i as add-on therapy to SoC in CLAD-BOS patients after single and double lung transplantation Despite the COVID-19 pandemic, BOSTON-1 and BOSTON-2 are ongoing, ensuring the integrity of data collection and safety of enrolled **V8 V7 V9** V10 patients **Roll-Over to** DISCLOSURES **BOSTON-3** The authors meet criteria for authorship as recommended by the International Committee of Medical Journal Editors (ICMJE) The authors received no direct compensation related to the development of the poster.

Patients with confirmed other causes for loss of lung function, such as acute infection, acute rejection, restrictive allograft syndrome (CLAD –RAS

- Patients with acute antibody-mediated rejection at Screening. In this context, clinically stable patients (as judged by the Investigator) with detectable levels of donor specific antibodies (DSA) at the Screening Visit are eligible for the
- Known hypersensitivity to L-CsA or to

