Corporate Presentation September 2022



Forward Looking Statements

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Our public communications, including this presentation, and SEC filings, may contain statements related to future, not past, events. These forward-looking statements are based upon current beliefs and expectations of Beyond Air's management and are subject to significant risks and uncertainties. These forward-looking statements often, but not always, may be identified by the use of words such as "believes," "estimates," "anticipates," "targets," "expects," "plans," "projects," "intends," "predicts," "may," "could," "might," "will," "should," "approximately," potential" or, in each case, their negative or other variations thereon or comparable terminology, although not all forward-looking statements contain these words. If underlying assumptions prove inaccurate or risks or uncertainties materialize, actual results may differ materially from those set forth in the forward-looking statements.

These forward-looking statements appear in a number of places throughout this presentation and include statements regarding our intentions, beliefs, projections, outlook, analyses or current expectations concerning, among other things, the patient market size and market adoption of our products by physicians and patients, the timing and cost of clinical trials for our products or whether such trials will be conducted at all, completion and receiving favorable results of clinical trials for our products, the development and approval of the use of nitric oxide for additional indications, FDA approval of, or other regulatory action with respect to, the timing, cost or other aspects of the commercial launch of our products and the commercial launch and future sales of our products or any other future products or product candidates. The extent to which the COVID-19 pandemic and global efforts to contain its spread will impact our operations, including the ability to conduct our preclinical studies and clinical trials or rely on our third-party manufacturing and supply chain, will depend on future developments, which are highly uncertain and cannot be predicted at this time, and include the duration, severity and scope of the pandemic and the actions taken to contain or treat the COVID-19 pandemic.

By their nature, forward-looking statements involve risks and uncertainties because they relate to events, competitive dynamics, and healthcare, regulatory and scientific developments and depend on the economic circumstances that may or may not occur in the future or may occur on longer or shorter timelines than anticipated or not at all. Although we believe that we have a reasonable basis for each forward-looking statement contained in this presentation, we caution you that forward-looking statements are not guarantees of future performance and that our actual results of operations, financial condition and liquidity, and the development of the industry in which we operate may differ materially from the forward looking statements contained in this presentation.





Harness the Power of Nitric Oxide



Ability to transition between hospital and untapped at-home market Creating Unlimited Access to On-Demand Nitric Oxide from Ambient Air

Nasdaq Listed: XAIR

Headquarters: Garden City, NY

Other Locations: Ireland, Israel



LungFit[®] Platform Elicits Paradigm Shift for Nitric Oxide Therapy



*Commonly referred to as persistent pulmonary hypertension of the newborn or PPHN.

+Caution: LungFit® PRO and LungFit® GO are investigational device, limited by federal (or United States) law to investigational use. High concentration iNO is defined as >80 ppm to <400 ppm. Low concentration is \leq 80 ppm.



Introducing the Patented Ionizer[™] Technology

The LungFit[®] PH is the first and only system with patented **lonizer technology** that generates nitric oxide using room air, enabling the delivery of **unlimited**, **on-demand nitric oxide** regardless of dose or flow.



Robust, Active Pipeline

	Therapeutic Area	Preclinical	Pilot	Pivotal	PMA	Commercial	Next Milestone ¹
LungFit [®] PH Hospital NICU Setting	Persistent pulmonary hypertension of the newborn (PPHN)			\rangle	\rangle		US FDA Approved CE Mark 2H22 ²
Low-concentration iNO (≤80 ppm) for pulmonary treatments	Cardiac surgery	\rangle		\rangle			US Submission 2H22
LungFit®PRO Hospital Setting	Viral Community-Acquired Pneumonia (VCAP), including COVID-19						Initiate US Study in
High-concentration iNO (80 to 400 ppm) for antimicrobial treatments	Bronchiolitis						Discussion with FDA
LungFit®GO At-Home Treatment High-concentration iNO (80 to 400 ppm) for antimicrobial treatments	Nontuberculous mycobacteria (NTM) lung infection						Full Dataset in 2H22 (home self-administration)
	Severe exacerbations due to lung infections in COPD patients						Pilot Study Start 2023
BEYOND CANCER [™] Next Level ImmuNO-oncology Next Level ImmuNO-oncology	Multiple solid tumors Visit <u>beyondcancer.com</u>						First in Human Trial Ongoing

1) All dates are based on projections and appropriate financing, anticipated first launch on a global basis pending appropriate regulatory approvals

6 2) Label expected to include cardiac surgery and PPHN



Large Market Opportunities



Beyond Air

7 9) Jinjuvadia, Chetna et al. . COPD. 2017;14(1):72-79.

10) Company Presentations and Regulatory Filings from Bristol-Myers Squibb, Merck, Roche, AstraZeneca, Pfizer, Regeneron; Sanofi 2011-2020.

The Critical Role Endogenous Nitric Oxide Plays in the Human Body



Nitric Oxide Has Multiple Mechanisms of Action



2) Saura, M., et al., An antiviral mechanism of nitric oxide: inhibition of a viral protease. Immunity, 1999. 10(1): p. 21-8

9

3) Akerström S et al. Nitric oxide inhibits the replication cycle of severe acute respiratory syndrome coronavirus. J Virol. 2005; 79(3):1966-9

4) Wink DA et al., Chemical biology of nitric oxide: Insights into regulatory, cytotoxic, and cytoprotective mechanisms of nitric oxide. Free Rad Biol Med 1998: (4-5): 434-56.

Nitric Oxide Plays a Major Role in the Immune System

Source of NO (cell type)	Category	Effector function
Macrophages, microglia, neutrophils, eosinophils, fibroblasts, endothelial cells, epithelial cells	Antimicrobial activity	Killing or reduced replication of infectious agents (viruses, bacteria, protozoa, fungi and helminths)
Macrophages, eosinophils	Anti-tumor activity	Killing or growth inhibition of tumor cells
Macrophages, microglia, astroglia, keratinocytes, mesangial cells	Tissue-damaging effect (immunopathology)	Necrosis or fibrosis of the parenchyma
Macrophages ('suppressor phenotype')	Anti-inflammatory — immunosuppressive effect	Immunoregulatory functions Inhibition of T and B cell proliferation, leukocyte recruitment (adhesion, extravasation, chemotaxis), Antibody production by CD5+B cells, autoreactive T and B cell diversification
Macrophages, T cells, endothelial cells, fibroblasts	Modulation of the production and function of cytokines, chemokines and growth factors	Up- and downregulation, e.g., of: IL-1, IL-6, IL-8, IL-10, IL-12, IL-18, IFN-γ, TNF TGF-β, G-CSF, M- CSF, VEGF, MIP-1α, MIP-2, MCP-1
Macrophages	T helper cell deviation	Induction and differentiation of TH1 cells Suppression of TH1 (and TH2) cell responses Suppression of tolerogenic T cell responses



Persistent Pulmonary Hypertension of the Newborn (PPHN)*

LungFit[®] PH offers significant advantages to hospitals

LungFit®PH

*PPHN is commonly used to refer to the condition treated with NO. The actual labelled indication is to improve oxygenation and reduce the need for extracorporeal membrane oxygenation in term and near-term (>34 weeks gestation) neonates with hypoxic respiratory failure associated with clinical or echocardiographic evidence of pulmonary hypertension in conjunction with ventilatory support and other appropriate agents.

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Nitric Oxide for PPHN and Cardiac Surgery



Inhaled Medical Gases: More to Breathe Than Oxygen, Michael A Gentile, Respiratory Care September 2011, 56 (9) 1341-1359
 Persistent Pulmonary Hypertension of the Newborn, Satyan Lakshminrusimha and Martin Keszler, NeoReviews December 2015, 16 (12) e680-e692;

3) Left ventricular heart failure and pulmonary hypertension, October 2015, European Heart Journal 37(12)

Current Nitric Oxide U.S. Market Dynamics

Established standard of care for 20+ years for pulmonary hypertension in the hospital setting (only PPHN on label)



LungFit[®] PH Receives FDA Approval in June 2022



1) Mallinckrodt Company Reports; Beyond Air estimates

2) American Academy of Pediatrics NICU Search

3) According to the CDC

4) Lakshminrusimha et al. 2015.

Incidence ~1.9 per 1,000 live births (range 0.4-6.8 per 1,000 births)⁴ ~7.5K Newborns in the US affected

by HRF every year

Lung-it[®]PH

Current Nitric Oxide U.S. Market Dynamics



2022 LungFit[®]PH receives FDA approval



1999 INOmax received FDA approval to treat hypoxic respiratory failure (HRF) in neonates

LungFit[®] PH: The Power to Transform iNO Care



Beyond Air Smart Filter vs. Cylinder

LungFit® PH: Revolutionary, Smart Design

Proprietary smart filter removes toxic nitrogen dioxide (NO₂)

Filters are a fraction of the cylinder size

- No disposal requirements
- · Easy to store, handle, and manage inventory

Smart filter (with RFID chip)

- · Measures time until filter change required
- Recognition LungFit® will not function without smart filter
 - Safety prevents NO₂ toxicity
 - Encryption prevents counterfeits
- Filter programs the system
 - Sets concentration and flow rate (not true for LungFit® PH)

Smart filter ensures hospital only charged for what is used

• Each filter lasts 12 hours regardless of dose or flow





Significant Advantages to Hospitals



Revolutionizing the Delivery of Nitric Oxide



An All-Inclusive Partnership

LungFlex™ 24/7 Partnership & Support

Exceeding your expectations with:

- All-inclusive contract—LungFit[®] PH Systems, backup systems, and accessories, creating budget certainty
- 24/7 live customer service and support (technical, clinical, commercial)
- LungFlex Rapid Replacement Program— Emergency deliveries within 6 hours*
- Convenient ordering for all components with first priority overnight deliveries
- On-demand, on-site clinical expertise and support

Comprehensive live on-demand training customized to the hospital's needs

*LungFlex Rapid Replacement Program response times are based on hospital locations. While every effort is made to accommodate emergency deliveries within 6 hours of request, some hospital locations may take longer.



LungFlex 24/7 Service and Support Line 1-855-LUNG-FLX or 1-855-586-4359



Harnessing the Power of Nitric Oxide

LungFit[®]PH

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Built on a Legacy of Innovation



LungFit[®] For Treating Lung Infections

Simple, safe and convenient

- · Allows for both home and hospital use
- Supplemental oxygen can be utilized through the system (hospital only)

Easy to Use

- Programmable by RFID on filter
- Convenient for all staff
- Self-administration for home use
- Usable with any electrical outlet 110/220V

Portable

• Only 20 lbs (home version may be lighter)

One system can treat multiple patients

- Easy to change breathing circuit
- One circuit per patient
- One filter per treatment

One Respiratory Therapist (RT) can operate multiple systems

- Insert filter and press "GO"
- Alarms monitor performance





High Concentration NO – Beyond Air Demonstrated Safety in Humans



- Beyond Air has 10+ years of experience with high concentration NO
- Concentrations as high as 250 ppm have been tested, with no SAE's
- Currently only 20 ppm NO approved by FDA
- Multiple animal studies in 2 species show intermittent dosing up to 400 ppm NO to be safe with no macroscopic or microscopic findings



Intermittent Dosing Key to Safe Administration of High Concentration NO

- Methemoglobin (MetHb) is a well-known biomarker for safety of NO; with the acceptable safety threshold <10%
- Methemoglobinemia is a condition where higher-thannormal levels of MetHb are found in the blood, causing too little oxygen to be delivered to the cells of the body
- An intermittent dosing regimen allows for high concentration NO to be administered without negative side effects, specifically addressing concerns of methemoglobinemia
- In the clinical study, MetHb levels followed a predictable pattern, rising during NO administration and falling back to normal, baseline levels shortly after the administration was stopped



Mean MetHb levels of 5 NO administrations (160 ppm every 4 hours) per day in 9 subjects for 14 days

Viral Community-Acquired Pneumonia in Hospitalized Patients

Nitric oxide has demonstrated antiviral and broad-spectrum antimicrobial activity



Viral Lung Disease Overview

Vaccines are not available for all causes of pneumonia



Viral Community-Acquired Pneumonia (VCAP)

- Influenza virus is the most common cause of viral pneumonia in adults¹
- Other viruses that cause viral pneumonia include¹: varicella-zoster virus, respiratory syncytial virus (RSV), human metapneumovirus, adenoviruses, picornaviruses, and coronaviruses
- Antibiotics are used for the bacterial causes of pneumonia, but are ineffective for viral causes²

Benefits of Nitric Oxide

- Nitric Oxide has broad-spectrum activity
 - Preclinical studies show high dose NO has antibacterial and antiviral properties³⁻⁴
 - Presented *in vitro* preclinical data at CHEST 2020 which support highconcentration NO has anti-coronavirus properties within hours
- Pulmonary vasodilatory properties
 - FDA/EMA approved for ~20 years

Bronchiolitis

- RSV is the most common cause of bronchiolitis in children⁵
- Usually affects children <2 years, with a peak in infants aged 3-6 months⁶
- Leading cause of infant hospitalizations, accounting for >120,000 hospitalizations with a direct cost of at least \$550 million each year⁶

Leading cause of childhood mortality



- 1) Cesario T., Viruses Associated With Pneumonia in Adults, Clinical Infectious Diseases, V. 55, I. 1, 1 July 2012, Pgs 107–113
- 2) American Thoracic Society- Top 20 Pneumonia Facts 2019 (here)
- 3) Saura, M., et al., An antiviral mechanism of nitric oxide: inhibition of a viral protease. Immunity, 1999. 10(1): p. 21-8
- 4) Wink DA et al., Chemical biology of nitric oxide.." Free Rad Biol Med 1998: (4-5): 434-56.
- 5) Piedimonte G, et al. Respiratory syncytial virus infection and bronchiolitis. Pediatr Rev. 2014; 35(12):519-30
- 6) Hasegawa et al. Trends in bronchiolitis hospitalizations in the United States, 2000-2009. Pediatrics 2013, 132(1):28-36.

Potential Nitric Oxide Market for Patients Hospitalized with Viral Respiratory Infections



- 1) According to the World Health Organization
- 2) Rudan et al. 2005.
- 3) According to the CDC
- 4) According to the national hospital ambulatory medical care survey in 2017
- 5) In 2019 according to UNICEF
- 6) Hall et al. Pediatrics Volume 132, Number 2, August 2013.
- 7) In 2019 according to the CDC

26



NO Tested in Three Bronchiolitis Pilot Trials

	Trial 1	Trial 2	Trial 3
Treatment groups	160 ppm NO + SST SST alone (control)	160 ppm NO + SST SST alone (control)	150 ppm NO + SST 85 ppm NO + SST SST alone (control)
Total Intent to Treat (ITT) Subjects Enrolled & Evaluated as the Safety Population	43	68	87
Study Treatment Protocol	Inhaled NO was given for Inhaled NO was given for 30 minutes, 5 times per day 30 minutes, 5 times per for up to 5 days for up to 5 days		Inhaled NO was given for 40 minutes, 4 times per day for up to 5 days
Primary objective	Safety	Efficacy (Length of Stay)	Efficacy (Time to Fit for Discharge)
Published or Presented	PEDIATRIC PULMONOLOGY OREANAL ARTICLE: RESPIRATORY INFECTIONS Nitric oxide inhalations in bronchiolitis: A pilot, randomized, double-blinded, controlled trial Adder Yales, Toxid Genetings, Toxing Acody, Idoal Golan-Tripte, Yaal Taivitsien, Shalom Ben-Sikimol, Ron Dagen, Adv. A Caddaar; First prainibilities 27 November 2017 https://doi.org/10.1002/pui.23055 [] Children: 1	SCIENTIFIC REPORTS Junitersearch Inhaled nitric oxide therapy in acute bronchiolitis: A multicenter randomized clinical trial	Annual Meeting

Next Steps:

Pivotal study on hold due to COVID-19 pandemic –

a study in VCAP may or may not precede a pivotal bronchiolitis study depending on pandemic conditions and conversations with FDA

NO Safe & Well Tolerated in Bronchiolitis Studies

Pooled Safety Results Presented at American Thoracic Society International Conference 2021

	SST (N=82)		85 ppm (N	NO + SST =32)	150 ppm (N	NO + SST =29)	160 ppm (N	NO + SST =55)	AII (M	V=198)
	Ν	%	Ν	%	Ν	%	Ν	%	Ν	%
Any AE	45	54.9%	20	62.5%	18	62.1%	25	45.5%	108	54.5%
Any SAE	10	12.2%	1	3.1%	3	10.3%	11	20.0%	25	12.6%

150 – 160 PPM NO treatment administered intermittently was generally safe and well tolerated across the three pilot trials, with the adverse event rates similar among treatment groups

Long-Term Safety Data Supports High Concentration NO in Bronchiolitis

Long-Term Safety Results Presented at Pediatric Academic Societies 2022 Meeting

Subjects re-hospitalized for bronchiolitis related outcomes

Treatment/ Control Group	Subjects Re- hospitalized (N)	Total Subjects (N)	Incidence Rate (95% CI)%	Patient Exposure Years*	Rate per 100 PEY (95% CI)
SST	6	39	15.39 (5.86 to 30.53)	143.0	4.20 (1.60 to 8.33)
85 PPM iNO + SST	1	24	4.17	38.0	2.63
150 PPM iNO + SST	1	20	5.00	32.4	3.09
160 PPM iNO +SST	2	18	11.11 (1.38 to 34.71)	90.6	2.21 (0.27 to 6.90)

*PEY=Patient Exposure Years, It is anticipated that the follow-up time when subjects completed the original studies to this current study will be different for different subjects. It is, therefore, necessary to calculate the patient year (PEY) which is the summation of the time (in years) from original study completion date to date of participation in the current study

- The multi-center study for longitudinal data collection was designed to evaluate the long-term effect of inhaled NO treatment in infants who participated in three pilot studies conducted between 2012-2020 and were given 85 – 160 PPM inhaled NO intermittently
- A total of 198 infants participated in the 3 studies, with 101 infants participating in long-term follow-up study
- Study concludes that the treatment of hospitalized infants with acute bronchiolitis by intermittent high dose NO show a favorable longterm safety profile and support further development of high concentration NO in this population

150 PPM NO is Minimum Therapeutic Dose

Data Presented at CHEST 2020 – Statistical Significance on both the Primary & Secondary Endpoint at 150 PPM

Third Bronchiolitis Pilot Study Results

	150 ppm vs. 85 ppm	150 ppm vs. SST	85 ppm vs. SST
Primary endpoint: Time to Fit-to-Discharge (FTD)			
Hazard Ratio	2.11	2.32	0.90
95% CI	1.03, 4.31	1.01, 5.33	0.44, 1.81
P-value	0.041	0.049	NS
Secondary Endpoint: Hospital Length of Stay (LOS)			
Hazard Ratio	2.01	2.28	0.77
95% CI	1.01, 3.99	1.03, 5.06	0.40, 1.48
P-value	0.046	0.043	NS



Viral Community-Acquired Pneumonia (VCAP) Pilot Study Design

Pilot Clinical Trial in Israel

- ✓ Commenced enrollment in November 2020
- Interim data presented at ATS 2021
- ✓ Additional data presented at ECCMID 2022
- Multicenter open label study of adult patients hospitalized with VCAP, including COVID-19
- Objective: establish 150 ppm NO is safe and tolerable in target patient population



150 PPM NO Evaluated in VCAP (including COVID-19) Pilot Study

Results Presented at the European Congress of Clinical Microbiology & Infectious Diseases in 2022

Intent to Treat Population: 35 subjects (16 iNO + SST vs 19 SST)

Demographics	SST	LungFit- 150 ppm NO+SST	All
Number	19	16	35
Age: Mean	53.2	50.5	51.9
Std	11.9	16.1	13.8
Min	20	23	20
Max	71	78	78
Female	11%	44%	26%

iNO treatment with LungFit® PRO was well tolerated

- No treatment was discontinued due to AE or discomfort
- No clinically significant differences were noted in respiratory rate, heart rate or blood pressure when compared between pre and end of inhalation
- MetHb levels were below 6.8% at all times (10% limit)
- NO₂ levels were below 4.4 ppm at all times (5 ppm limit)

Baseline Characteristics	SST	LungFit- 150 ppm NO+SST
O2 required %	68.4	62.5
Cardiac disorders %	10.5	12.5
Metabolic disorders %	47.4	43.8
Respiratory disorders %	21.1	12.5
Vascular disorders %	21.1	50.0
Adverse Events	SST	LungFit- 150 ppm NO+SST
Adverse Events Any AE	SST 9 (47%)	LungFit- 150 ppm NO+SST 13 (81%)
Adverse Events Any AE Any AE drug/device related*	SST 9 (47%) 0	LungFit- 150 ppm NO+SST 13 (81%) 0
Adverse Events Any AE Any AE drug/device related* Any SAE	SST 9 (47%) 0 0	LungFit- 150 ppm NO+SST 13 (81%) 0 2 (13%)
Adverse Events Any AE Any AE drug/device related* Any SAE Any SAE drug/device related*	SST 9 (47%) 0 0 0 0	LungFit- 150 ppm NO+SST 13 (81%) 0 2 (13%) 0
Adverse Events Any AE Any AE drug/device related* Any SAE Any SAE drug/device related* Any AE (moderate or severe)	SST 9 (47%) 0 0 0 3 (16%)	LungFit- 150 ppm NO+SST 13 (81%) 0 2 (13%) 0 3 (19%)

*including possibly and probably related

VCAP (including COVID-19) Efficacy Data

Efficacy Data show Strong Trends in Favor of NO (97% of subjects were Covid-19)

Intent to Treat Population: 35 subjects (16 iNO + SST vs 19 SST)



Source: ECCMID Poster Presentation; Based on Cox proportional hazard model

Nontuberculous Mycobacteria

Expanding Nitric Oxide into the Home Market for Lung Infections



Nontuberculous Mycobacteria (NTM) Overview

• Who is at risk?

Immunocompromised people are at a greater risk for NTM



¹⁾ Donohue et al. Environ Sci Technol. 2015;49(10):6127-6133. 2) Hall-Stoodley et al. Biofilm formation by the rapidly growing mycobacterial species Mycobacterium fortuitum. FEMS Microbiol Lett. 1998;168(1):77-84.

How is NTM acquired?

NTM is commonly found in water sources, with warmer climates having higher infection rates



Gulf States account for

in the United States¹

70% of annual NTM cases

Patient to patient



species of bacteria



US study across 25 states showed that NTM bacteria were found in nearly 8 out of 10 water samples²



Bacteria live in soil from parks, gardens, and environment



35

Home Market: NTM Market Dynamics

LungFit[®]GO



1) Kotilainen, H. et al. "Clinical Findings in Relation to Mortality in Non-Tuberculous Mycobacterial Infections..."European Journal of Clinical Microbiology & Infectious Diseases 34.9 (2015)

2) Strollo et al. The Burden of Pulmonary Nontuberculous Mycobacterial. Pub 27-July-2015

3) Pyarali FF, Schweitzer M, Bagley V, et al. Increasing non-tuberculous mycobacteria infections in veterans with COPD and association with increased risk of mortality. Front Med (Lausanne). 2018;5:311.

4) Data presented at ATS 2017 (Derek Low et al, Medical University of South Carolina)

36

Current NTM Market Dynamics

Targeting Refractory Mycobacterium avium complex (MAC) & M. abscesses NTM Patients



1) Winthrop et al. Incidence and prevalence of nontuberculous mycobacterial lung disease in a large U.S. managed care health plan, 2008-2015. Ann Am Thorac Soc, 17 (2020), pp. 178-185

2) Ringshausen et al. Prevalence of Nontuberculous Mycobacterial Pulmonary Disease, Germany, 2009-2014. Emerg Infect Dis. 2016;22(6):1102-1105.

3) Izumi et al. Epidemiology of Adults and Children Treated for Nontuberculous Mycobacterial Pulmonary Disease in Japan. Ann Am Thorac Soc. 2019 Mar;16(3):341-347.

4) Diel R et al. High mortality in patients with MAC lung disease: a systematic review. BMC Infect Dis. 2018;18(1):206. Published 2018 May 3

5) According to the Cystic Fibrosis Foundation

37

Pilot Study in CF Patients with NTM Lung Infections Demonstrates Safety and Efficacy

Single arm study with 160 ppm NO showed a reduction in bacterial load and improvements in quality of life Data Published in the Journal of Cystic Fibrosis (Bentur et al., 2019)



- 9 CF patients with refractory MABSC were treated at 3 centers in Israel with NO added to background antibiotic therapy
 - 160 ppm NO was given via mask for 30 min 5x/day for 14 days and 3x/day for 7 days
 - Primary endpoint of safety was met, with no NO-related serious adverse events (SAEs) observed
 - Bacterial load, as measured by qPCR showed a 65% reduction at day 81 versus baseline
 - One patient was culture negative at Day 51 and Day 81, two others had one negative culture
 - Quality-of-Life data showed positive trends on relevant questions
- 4 patients treated under compassionate use experienced similar results
 - 1 treated at NIH with LungFit[®], 1 treated safely with 250 ppm NO, 1 culture conversion

Pilot LungFit[®] NTM Study Protocol Summary

Pilot Clinical Trial In Australia



- Received grant for up to \$2.17 million from the Cystic Fibrosis Foundation to help fund pilot study
- ✓ Interim results presented at the American Thoracic Society (ATS) 2022 International Conference in May 2022
- 12-week, single-arm, multicenter study enrolling up to 20 adult Cystic Fibrosis (CF) or non-CF bronchiectasis patients with refractory NTM lung infections including *Mycobacterium avium* complex (MAC) and *Mycobacterium abscessus* complex (MABSC)



Positive Interim Results for LungFit[®] GO Pilot Study Presented at ATS 2022: NO Self-Administered At-Home

Baseline Demographics				
	Ν	15		
	Mean	62.1		
Age (vrs)	Std	15		
/ go (yro)	Min	22		
	Max	82		
Gender	Male	3		

Adverse Events (ITT** Population) = 15	Ν	%
Any AE	14	93.3
Any AE related to study treatment *	9	60.0
Any AE related to study treatment classified as Severe *	0	0
Any SAE	6	40.0
Any Treatment Emergent SAE occurring during treatment period	2	13.3
Any SAE related to study treatment *	1	6.7

*Including possibly, probably, and definitely related **Intention to treat

- All subjects completed the in-hospital phase and were titrated to a 250 ppm NO regimen at home
- A total of 300 inhalations were administered in-hospital, with mean SpMetHb of 4.4% and NO₂ of 3.5 ppm at the end of inhalation
- To date, a total of 2323 inhalations were self administered at home with no treatment related AE discontinuations reported and overall high treatment compliance

Positive Interim Results for LungFit[®] GO Pilot Study Presented at ATS 2022: NO Self-Administered At-Home

Quality of Life (QoL-B) Overall QoL improvement shown in Percentage of Subjects Improved/Remained Unchanged majority of categories at Day 85*** unchanged ■ improved Latest FDA draft guidance for NTM-100 of Subjects pulmonary disease caused by MAC is 90 80 "To support approval, FDA 70 expects that drugs will provide 60 benefit on a clinically meaningful 50 Percentage endpoint" 40 30 "Primary efficacy endpoints should 20 be based on clinical outcome 10 assessments, such as a PRO 0 Digestive Symptoms tons problems under the stimp of the symptoms and symptom instrument assessing symptoms" Vitality Expect further efficacy data/analysis • later in 2022 QoL **Bacterial** load **Physical function** ***Calculated only for subjects completing treatment period

How Big is the Home Market for Severe Lung Infections?



- ...is the largest at-risk population for recurrent and opportunistic lung infections
- There are an estimated 30m people in the US suffering from COPD¹ with 10% considered severe²
- 1,075,575 estimated acute COPD exacerbation-related hospitalizations in 2010
- Average COPD exacerbation hospital LOS was 6 days in 2010
- \$38,455 cost per hospitalization in 2010 translates to >\$41b in cost

- After hospitalization varies between 16% and 19% in the 3 months following hospitalization, between 23% and 43% at 1 yr and is 55–60% at 5 yrs⁴.
- In the ECLIPSE⁵ study (Hurst et al. NEJM 2010), a 3 year observation of 1,679 moderate to severe COPD patients (GOLD 2,3 & 4)
 - 77% of patients had at least one exacerbation during the observation period
 - 47% of patients had <u>></u>2 exacerbations in at least one of the three study years
 - 30% of patients had ≥1 exacerbation in each of the three study years
 - 12% of patients had <u>></u>2 exacerbations in each of the three study years

1) COPD Foundation

- 2) Mannino and Braman: Epidemiology and Economics of COPD. American Thoracic Society. Volume 4. 2007
- 3) Jinjuvadia et al. Journal of Chronic Obstructive Pulmonary Disease 2017;
- 4) Raherison C and Girodet PO. Epidemiology of COPD. Eur Respir Rev. 2009;18(114):213-221;
- 5) Evaluation of COPD Longitudinally to Identify Predictive Surrogate Endpoints

BEYOND CANCER[™]

Next level immuNO-oncology

Nitric Oxide for Solid Tumors

An Affiliate of Beyond Air



Nitric Oxide is a Powerful Anti-Cancer Agent



NO has shown anticancer properties at ultra-high concentrations by activating the immune system

Our data suggest that our innovative gaseous NO-based treatment may treat solid tumors locally and their distant metastases systemically via stimulation of an anti-tumor immune response

Hypothesis: Exogenous ultra-high concentration NO (UNO; >10,000 ppm) administered directly to a solid tumor may cause local cell death resulting in systemic exposure to tumor antigens. Tumor antigens may trigger a systemic immune response, thereby creating a memory immune bank that will recognize and attack subsequent primary tumor regrowth as well as distal metastases.



Project UNO Will Target Patients with Solid Tumors



Solid Tumors represent approximately 90% of adult human cancers¹, accounting for approximately 1.4 million annual new cases of most common cancer types in the United States³

Metastatic Disease is responsible for 90% of solid tumor deaths²



Cooper GM. The Cell: A Molecular Approach. 2nd edition. Sunderland (MA): Sinauer Associates; 2000. The Development and Causes of Cancer. Available from: https://www.ncbi.nlm.nih.gov/books/NBK9963/ 45

2) Fontebasso Y, Dubinett SM. Drug Development for Metastasis Prevention. Crit Rev Oncog. 2015;20(5-6):449-473. doi:10.1615/CritRevOncog.v20.i5-6.150

3) According to the National Cancer Institute: https://www.cancer.gov/types/common-cancers

>\$30 Billion Checkpoint Inhibitor Market in 2021 and Growing

Company	Drug Name	First FDA Approval	2021 Revenue
Bristol-Myers Squibb	Yervoy	March 2011	\$2.0 Billion
Merck	Keytruda	Sept 2014	\$17.2 Billion
Bristol-Myers Squibb	Opdivo	Dec 2014	\$7.5 Billion
Roche	Tecentriq	May 2016	\$3.0 Billion
AstraZeneca	Imfinzi	May 2017	\$2.4 Billion

Source: Company Presentations and Regulatory Filings from Bristol-Myers Squibb , Merck , Roche, AstraZeneca 2011-2021

XTT cell proliferation-based assay

Human ovarian and pancreatic cancer cell lines were exposed to gaseous nitric oxide at 10,000 ppm - 100,000 ppm NO for 10 seconds to 10 minutes. Cell viability was assessed by XTT assay. Less than 10% of both cell lines are viable after 1 minute of exposure to 25,000pm NO.

■ Non treatment ■ NO 10,000 ppm ■ NO 25,000 ppm ■ NO 50,000 ppm ■ NO 100,000 ppm

■ Non treatment ■ NO 10,000 ppm ■ NO 25,000 ppm ■ NO 50,000 ppm ■ NO 100,000 ppm

One-way ANOVA and Dunnett's multiple comparison test, compared to non treatment, * p<0.05, ** p<0.01, *** p<0.001, **** p<0.001

Short exposure of ultra-high concentration gNO limits cell viability in B E Y O N D C A N C E R mouse colon and breast cancer cell lines

Clonogenic assay D CT26 (mouse colon cancer cells) 120% **** *** 0% 10 sec 1 min 2.5 m in Exposure time 4T1 (mouse breast cancer cells) F 120% 100%

One-way ANOVA and Dunnett's multiple comparison test, compared to non treatment, * p<0.05, ** p<0.01, *** p<0.001, **** p<0.001

Effects of UNO on CT26 Challenge Tumors In Vivo

Dose-dependent effects of primary tumor treatment on survival was observed in CT26 tumor-bearing mice treated with 20,000 or 50,000 ppm NO

Challenge assay: CT26 study mice were treated with either 20,000 or 50,000 ppm NO for 5 minutes. Naïve mice inoculated with the same cancer cells served as an internal control. Up to 21 days post NO treatment, all mice were re-inoculated with colon cancer cells (CT26 cells) as a challenge tumor and survival was monitored.

 At day 45, 25% of naïve mice, 73% of 20,000 ppm NO mice and 100% of 50,000 ppm NO mice were alive

Results:

Mechanism of Action Summary: Estimated Timeline Day 14+

Approximately 14+ days after treatment the adaptive and innate immune system is activated with increased levels of Tcells and B-cells circulating in the blood stream following ultra-high concentration gNO treatment.

Blood B-cells

Financial and Patent Information

Financial Overview

As of June 30, 2022

Patent Information

20 issued patents expiring through 2033

> 10 pending patents, if issued, may extend the last expiration through 2042

Beyond Air believes that its patent portfolio is strong and broad:

- The nitric oxide generator
- The breathing circuit
- Nitric oxide concentration
- Nitric oxide action in the body
- Nitric oxide dosing
- NO₂ filter
- Method of Use
- Cancer
- Coronavirus

Achievements and Upcoming Milestones

		1H22	2H22	1H23	2H23
LungFit [®] PH	Persistent Pulmonary Hypertension of the Newborn (PPHN) worldwide, and cardiac surgery outside US where indicated	US Commercial Approval & Launch	CE Mark US Cardiac PMA Supplement Submission	US Cardiac Label decision Ex-US Partnership Ex-US Launches Continues	
PRO	Viral Community-Acquired Pneumonia (VCAP), including COVID-19	VCAP data presentation at the 32 nd European			
LungFit	Bronchiolitis	Microbiology and Infectious Disease in April 2022"			Pandemic Permitting)
JFit°GO	Nontuberculous mycobacteria (NTM) Lung Infection (home self-administration)	Efficacy & Safety Data Presentation at ATS in May 2022	Complete NTM Pilot Study	Discuss next steps with FDA	Discuss next steps with FDA
Lung	Severe Exacerbations due to Lung Infections in COPD Patients				Potential Initiation of Study in COPD Patients
		Initiate Human Study	Announce Initial Human Data	Potential IPO	Potential IPO
	Next Level ImmuNO-oncology Multiple Solid Tumors	Presentation at AACR in April 2022	Preclinical Combination Data Presentation	Clinical and Preclinical Data Presentations	Initiate Additional Phase I Studies in Various Tumors

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