Synlogic

DESIGNED FOR LIFE

July 2019 Corporate Presentation



Forward Looking Statements

This presentation contains "forward-looking statements" that involve substantial risks and uncertainties for purposes of the safe harbor provided by the Private Securities Litigation Reform Act of 1995. All statements, other than statements of historical facts, included in this presentation regarding strategy, future operations, future financial position, future revenue, projected expenses, prospects, plans and objectives of management are forward-looking statements. In addition, when or if used in this presentation, the words "may," "could," "should," "anticipate," "believe," "estimate," "expect," "intend," "plan," "predict" and similar expressions and their variants may identify forward-looking statements. Examples of forward-looking statements include, but are not limited to, the approach we are taking to discover and develop novel therapeutics using synthetic biology; statements regarding the potential of our platform to develop therapeutics to address a wide range of diseases, including: inborn errors of metabolism, liver disease, inflammatory and immune disorders, and cancer; the future clinical development of Synthetic Biotic medicines; the potential of our technology to treat hyperammonemia and phenylketonuria; the expected timing of our anticipated clinical trial initiations; the benefit of orphan drug and fast track status; the adequacy of our capital to support our future operations and our ability to successfully initiate and complete clinical trials; the results of our collaborations; and the difficulty in predicting the time and cost of development of our product candidates. Actual results could differ materially from those contained in any forward-looking statement as a result of various factors, including, without limitation: the uncertainties inherent in the preclinical development process; our ability to protect our intellectual property rights; and legislative, regulatory, political and economic developments, as well as those risks identified under the heading "Risk Factors" in our filings with the SEC. The foregoing review of important factors that could cause actual events to differ from expectations should not be construed as exhaustive and should be read in conjunction with statements that are included herein and elsewhere, including the risk factors included in our Annual Report on Form 10-K filed with the SEC on May 9, 2019. The forward-looking statements contained in this presentation reflect our current views with respect to future events. We anticipate that subsequent events and developments will cause our views to change. However, while we may elect to update these forwardlooking statements in the future, we specifically disclaim any obligation to do so. These forward-looking statements should not be relied upon as representing our view as of any date subsequent to the date hereof.





Synthetic Biotic™ Medicines A Novel Class of Engineered Living Medicines

SYNTHETIC

- Designed genetic circuits to execute biological functions
- Degradation of disease-causing metabolites
- Production of therapeutic molecules

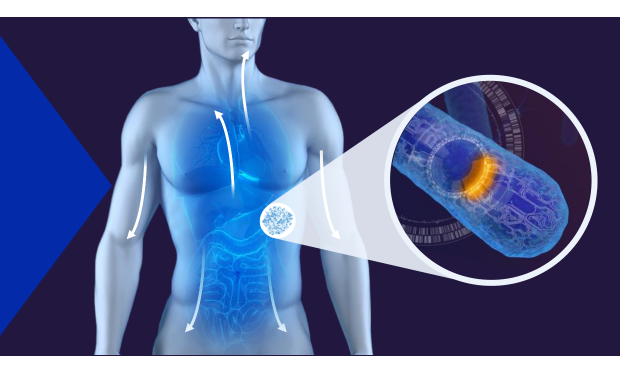
BIOTIC

- Bacterial chassis
- Non-pathogenic
- Amenable to genetic manipulation

PATHWAYS, COMBINATIONS, BIOMARKERS

PROGRAMMABLE POTENCY AND CONTROL

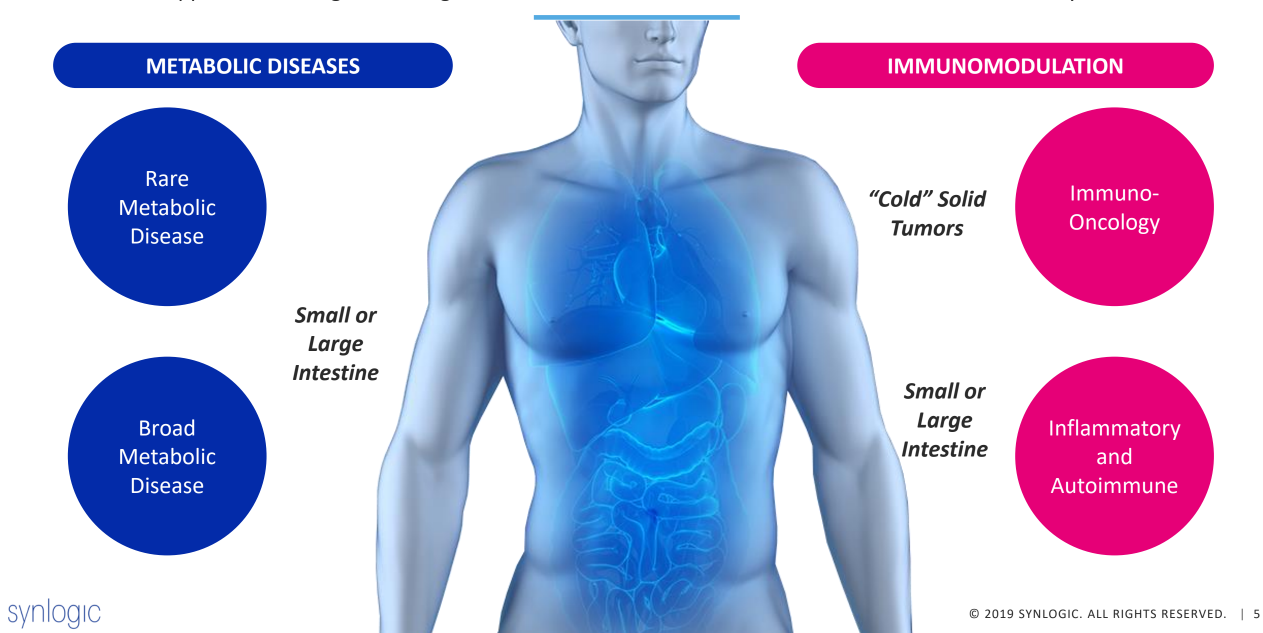
LOCAL ACTIVITY, REDUCED SYSTEMIC TOXICITY





Synthetic Biotic Portfolio: Breadth and Potential

Initial Applications Designed to Target Different Sites of Action in Metabolic and Immunomodulatory Diseases



Investing in Development of a Robust Pipeline for a Range of Diseases

| | Research IND-Enabling Phase 1 Phase 2 |
|--|---------------------------------------|
| Hyperammonemia – Urea Cycle Disorder | SYNB1020 |
| Phenylketonuria | SYNB1618 |
| Additional Rare Metabolic Diseases | |
| Hyperammonemia – Hepatic Encephalopathy (HE) | SYNB1020 |
| Inflammatory Bowel Disease | abbyie |
| Immuno-Oncology Solid Tumors | SYNB1891 |
| Additional Oncology Applications | |

Rare Metabolic Diseases

Broad Metabolic Disease

Immunomodulation



SYNB1020 for Hyperammonemia Indications

Characterized by Systemic Ammonia Accumulation

HEPATIC ENCEPHALOPATHY (HE)

Neuropsychiatric complication in patients with end-stage liver disease (cirrhosis)

- Liver dysfunction leads to ammonia accumulation
- Toxic to brain, leading to HE crisis & hospitalization

Patients:

- 165,000 diagnosed overt patients in US
- Up to 70% of patients with cirrhosis characterized as covert (subclinical)

Treatment:

- Lactulose: laxative with significant side effects
- Rifaximin: reduction in overt HE recurrence

Target Profile to Address Unmet Need:

- Reduce episodes of hospitalization
- Improve cognitive outcomes, Quality of Life

UREA CYCLE DISORDERS (UCD)

Genetic defects in Urea Cycle

- Deficiency in one of the six enzymes
- Nitrogen accumulates as toxic ammonia leading to metabolic crisis

Patients:

• ~2,000 diagnosed in US; similar in EU

Treatment:

- Ammonia scavengers: Buphenyl® (sodium phenylbutyrate), Ravicti® (glycerol phenylbuterate)
- Low protein diet with amino acid supplements

Target Profile to Address Unmet Need:

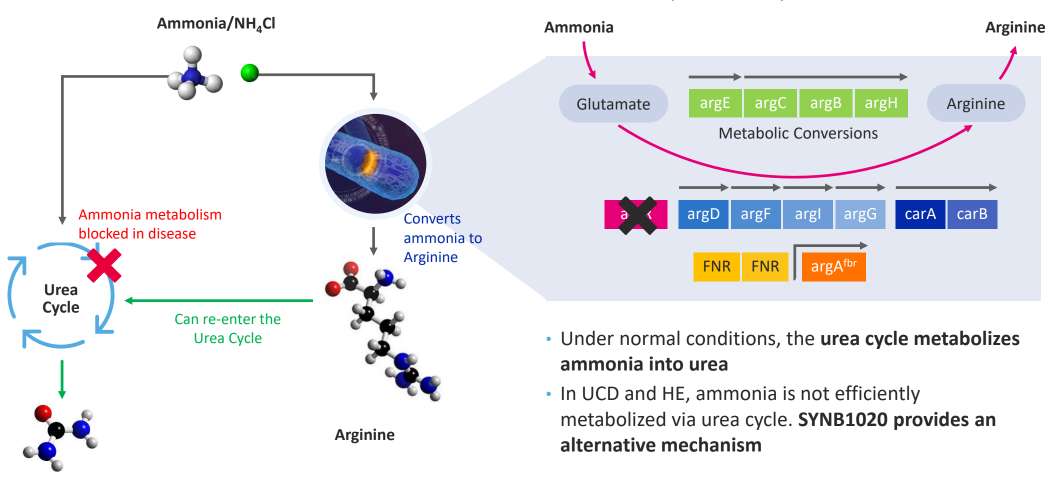
- Maintain blood ammonia in normal range, avoid crisis
- Protein liberalization: 50-100% more per day
- Oral administration



SYNB1020 Mechanism of Action:

Conversion of Toxic Ammonia into Beneficial Arginine for the Treatment of UCD and HE

Engineered Probiotic Bacteria: E. coli Nissle Components of Synthetic Genetic Circuit



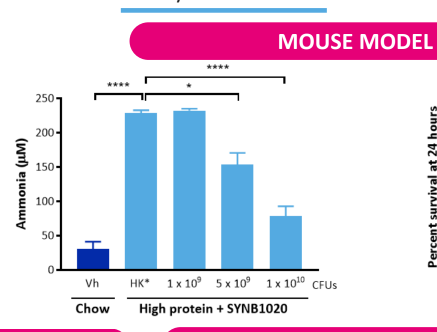


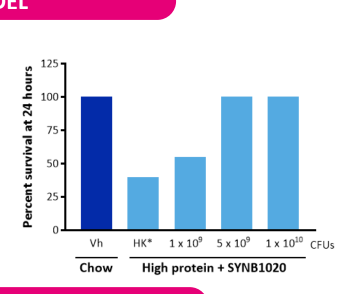
Urea

SYNB1020 data recently published in Science Translational Medicine

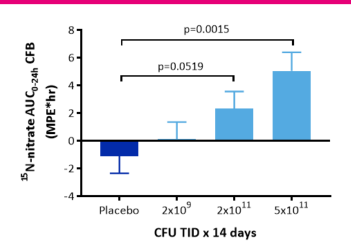
In vivo data in mouse models and healthy volunteers demonstrate mechanism of action



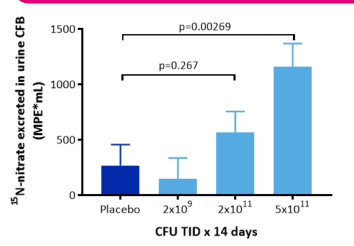




PLASMA NITRATE



URINARY NITRATE



SYNB1020 Clinical Data in Healthy Volunteers

Dose-dependent Increase in SYNB1020 in Feces, Clearance on Cessation of Dosing

DOSE-DEPENDENT INCREASE IN FECES

 $6x10^{11}$

Dose of SYNB1020 per day

Dosing period = 14 days Samples collected daily

6x10⁹

30000 -

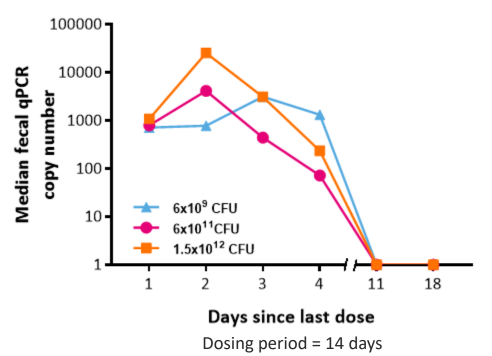
20000 -

10000 -

Fecal qPCR copy number (Mean within subject)

1.5x10¹²

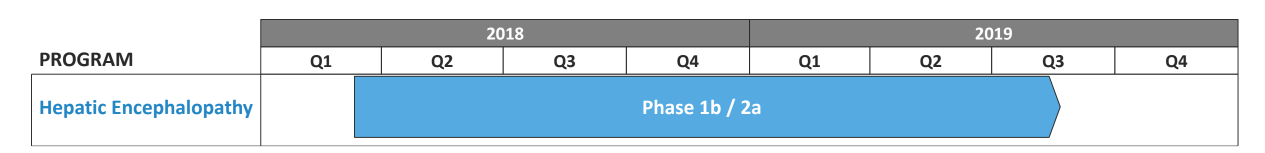
CLEARANCE





SYNB1020 Clinical Development

Hepatic Encephalopathy Phase 1b/2a in Patients with Cirrhosis and Elevated Ammonia



Hepatic Encephalopathy Clinical Trial

- Randomized, double-blind placebo-controlled study ongoing at multiple sites in the US
- Primary outcome: establish safety/tolerability in patients with cirrhosis and elevated ammonia
- Secondary outcome: reduction of ammonia

Urea Cycle Disorders

(Plans to continue development in UCD dependent on data from Ph 1b/2a HE study)





SYNB1618 for Phenylketonuria (PKU)

Goal: Managing Plasma Phe Levels

PKU is a rare inherited amino acid metabolism disorder

- Causes build up of amino acid phenylalanine (Phe) in the body
- Today, less than half of adults are at or below target Phe levels of 120-360 μmol / L
- If left untreated, symptoms include cognitive impairment, convulsions, behavioral problems, skin rash

Patients:

16,500 diagnosed in US, similar in EU5

Treatment:

- Phenylalanine is found in all proteins therefore low protein diet is followed (no meat, dairy, nuts, eggs)
- KUVAN® (sapropterin dihydrochloride): PAH cofactor. 20-40% of patients are responders
- Palynziq™ (pegvaliase-pqpz): injectable, pegylated, bacterial enzyme (phenylalanine ammonia-lyase or PAL) for treatment of adult patients

Target Profile to Address Unmet Need:

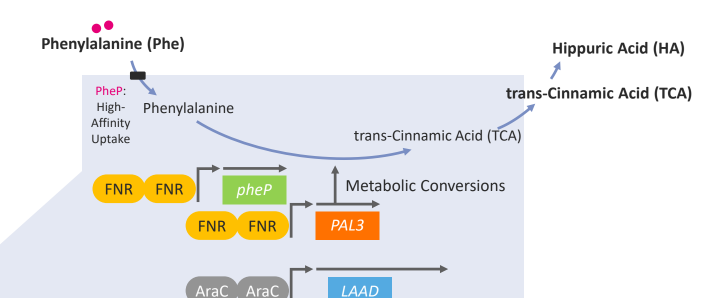
- Manage Phe below target levels to prevent irreversible cognitive damage
- Increase natural protein intake: classic PKU patients' natural protein intake is typically less than 10g
- Oral dosing without systemic toxicity



SYNB1618 Mechanism of Action

Amino acids from dietary proteins (absorption and recirculation) **Healthy** Phe Impaired PAH Phenylalanine Hydroxylase (PAH) converts Accumulation of Phe into Tyrosine Phe to toxic levels **SYNB1618** Tyrosine **Manage Phe levels**

Engineered Probiotic Bacteria: E. coli Nissle Components of Synthetic Genetic Circuit



When Phe is not efficiently metabolized (PKU) **SYNB1618** provides an alternative mechanism

- PAL3: produces TCA which is converted to HA in the liver and is excreted in urine
- LAAD: produces phenylpyruvate (PP)

Phenylalanine (Phe)



Phenylpyruvate (PP)

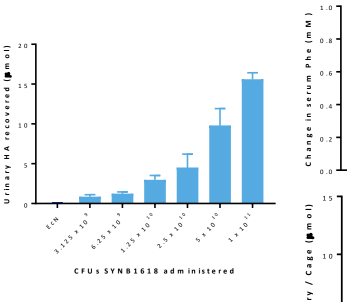
SYNB1618 Preclinical Characterization

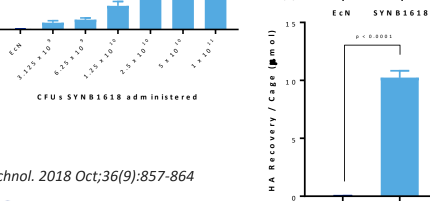
Biomarkers Demonstrate Activity of SYNB1618 in Mouse Model of PKU and Healthy NHPs



Development of synthetic live bacterial therapeutic for the human metabolic disease phenylketonuria Vincent M Isabella et al, Synlogic, Inc.

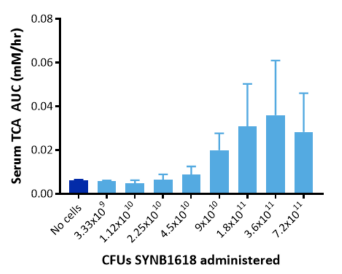
IN VIVO EFFICACY IN (PKU) PAHenu2/enu2 MOUSE

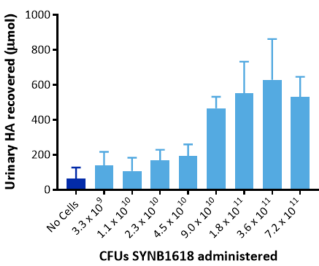




SYN B 1 6 1 8

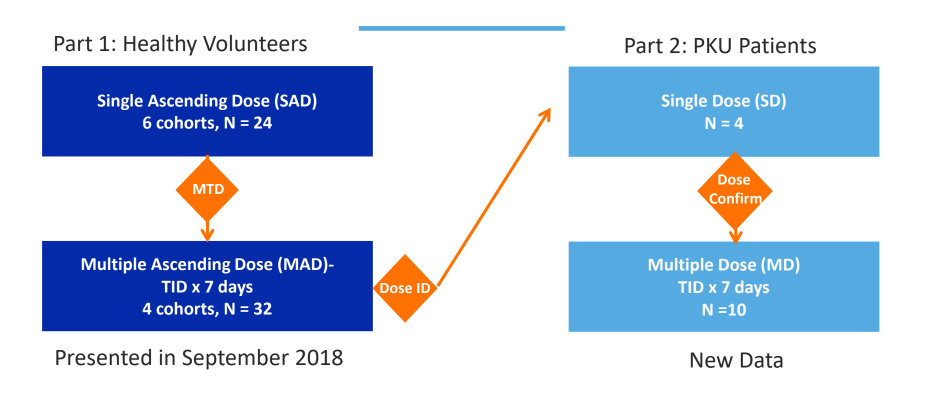
DOSE RESPONSE IN HEALTHY NHPs





Nat. Biotechnol. 2018 Oct;36(9):857-864

SYNB1618 Phase 1/2a Study Design



PKU Clinical Trial Design

- Randomized, double-blind placebo-controlled study at multiple sites in the US
- Primary outcome: establish safety/tolerability following single and multiple doses in HV and PKU patients
- Secondary outcome: SYNB1618 kinetics in feces
- Exploratory: change from baseline in plasma and urinary biomarkers of Phe metabolism



SYNB1618 in the Clinic: Safety

Phase 1/2a SAD/MAD Study Demonstrates Safety and Clearance in Healthy Volunteers and PKU Patients

56 healthy volunteers, 14 PKU patients

Received at least one dose of SYNB1618 or placebo

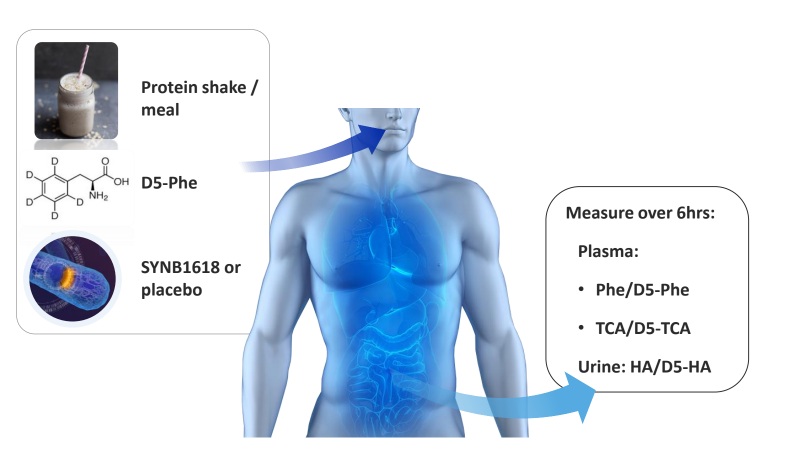
Adults Age range: 18-62 yrs old

- ✓ There were no treatment-related serious adverse events, no systemic toxicity or infections
- ✓ Treatment-emergent adverse events were either mild or moderate in severity, and reversible. Most adverse events were GI-related
- ✓ Single dose MTD in healthy volunteers was defined as $2x10^{11}$ CFU. Doses above this level were associated with dose-limiting GI adverse events
- \checkmark Based on pharmacodynamic data and tolerability profile, a dose of $7x10^{10}$ CFU was identified for the second part of the study in PKU patients
- ✓ Dose of 7x10¹0 CFU TID over seven days was well-tolerated in PKU patients. There were no discontinuations.
- ✓ All subjects cleared the bacteria (one PKU patient in follow-up). There was no evidence of colonization, and no subject required antibiotics

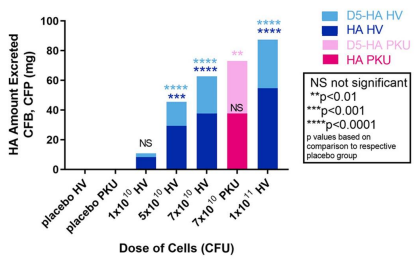


SYNB1618 in the Clinic: Activity

Statistically Significant and Equivalent Activity of SYNB1618 in Healthy Volunteers and Patients

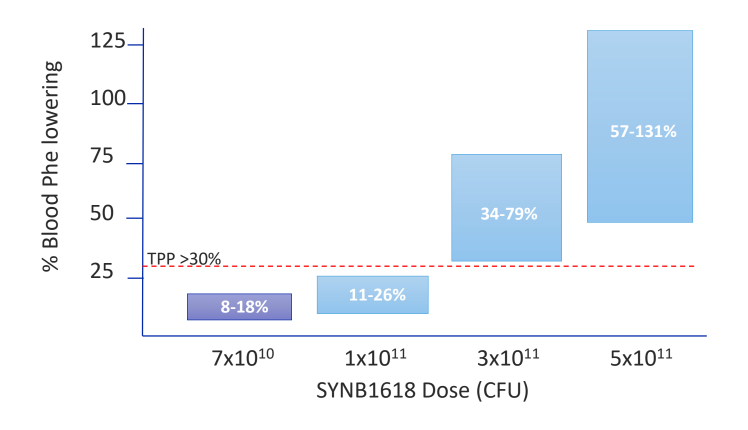


MD URINARY HA AND D5-HA



HA=hippurate, D5-HA= labeled HA, CFB=change from baseline, CFP=change from placebo HV=healthy volunteer PKU=phenylketonuria patient

Modeling: Potential For Phe Reduction in PKU Patients

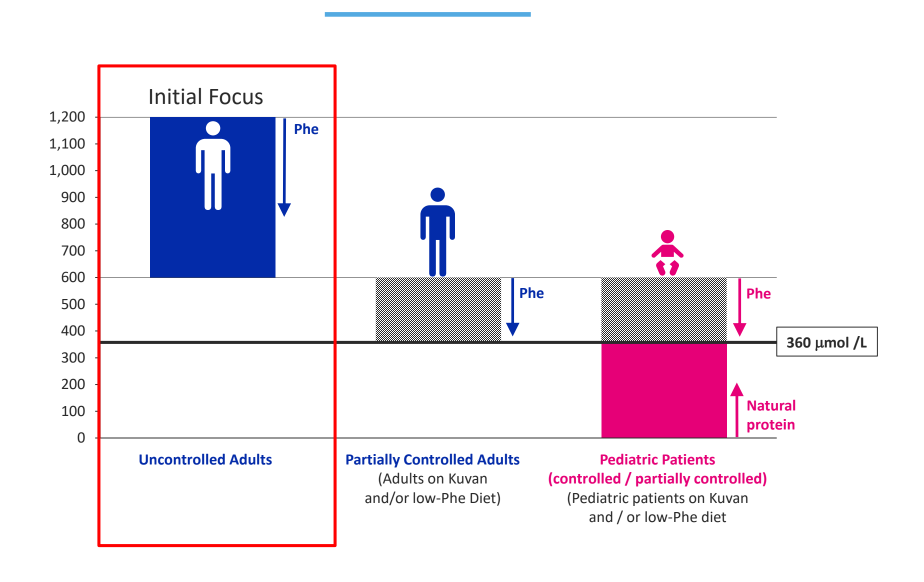


Ranges represent

- Low: PAL mechanism only (conservative)
- High: PAL + LAAD activity (estimates maximum with both pathways)



SYNB1618 Potential to Address Unmet Need Across Patient Groups





Upcoming Milestones and Path Forward

Established new solid formulation and manufacturing process



Completed EPO1 interactions with FDA to align on program plans (clinical, manufacturing, toxicology)



Completed Phase 1/2a study (healthy volunteers and PKU patients)



Initiate bridging study with solid formulation in Q3 2019

Phase 2 study in PKU patients to assess Phe lowering to start in 1H 2020





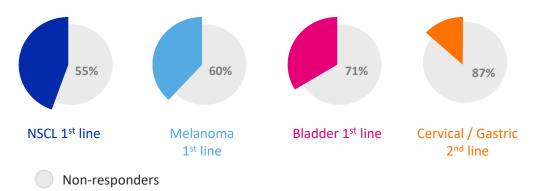
Synlogic Vision for Immuno-Oncology

Expand the Benefits of Immunotherapy Broadly Across Tumor Types

CHECKPOINT INHIBITORS HAVE TREATMENT FAILURES

For indications where immune checkpoint inhibitors are indicated, 55-87% of patients fail to respond

Failure Rates for Select FDA Approved CPI Monotherapy



Other tumors, where CPIs are not indicated, show littleto-no response to checkpoint inhibitors

Bacteria Recognized as Earliest Immunotherapy

Nature often gives us hints to her profoundest secrets, and it is possible that she has given us a hint in which, if we will but follow, may lead us on to the solution of this difficult problem.



DR. WILLIAM B. COLEY
IMMUNO-ONCOLOGY PIONEER

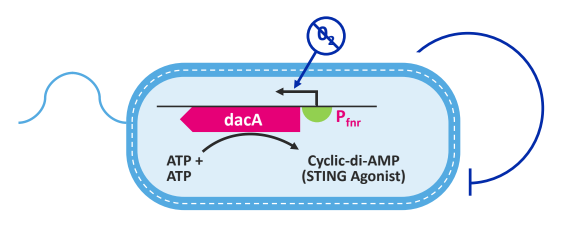
Enable broad response and remission through engagement of multiple immunomodulatory pathways to enhance tumor inflammation and promote robust T cell responses



Dual Innate Immune Activator: Synthetic Biotic Medicine Producing STING Agonist (SYNB1891)

- Synthetic biology applied to confer activities for efficacy and control for safety
- Designed as a dual innate immune activator: combined benefit of bacterial chassis and STING agonist
- The dacA gene is integrated into genome under the control of inducible promoter (P_{fnr}) to produce c-di-AMP (CDA)
- Dual biosafety feature via auxotrophies no proliferation in tumor, systemic circulation or environment
- Learnings inform future combinations

ANAEROBIC ENVIRONMENT



Auxotrophies

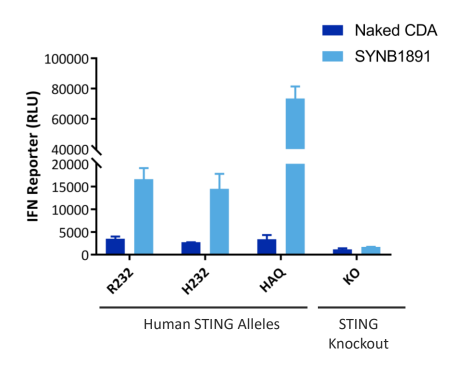
- Diaminopimelic acid (DAP)
- Thymidine



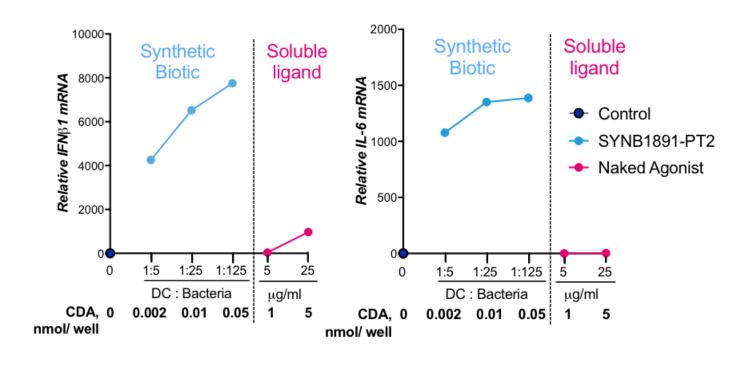
SYNB1891 In Vitro Characterization

Interferon Production Across Multiple Human STING Alleles – Activity Greater than Naked STING Agonist

REPORTER HUMAN MONOCYTIC LINE



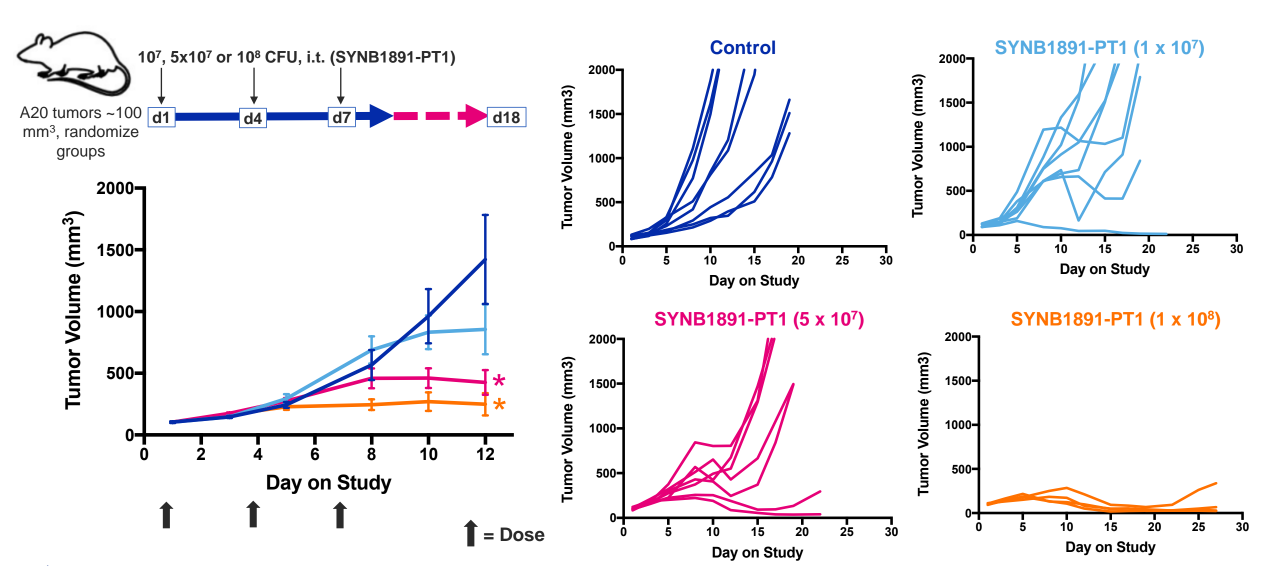
HUMAN PRIMARY DENDRITIC CELLS





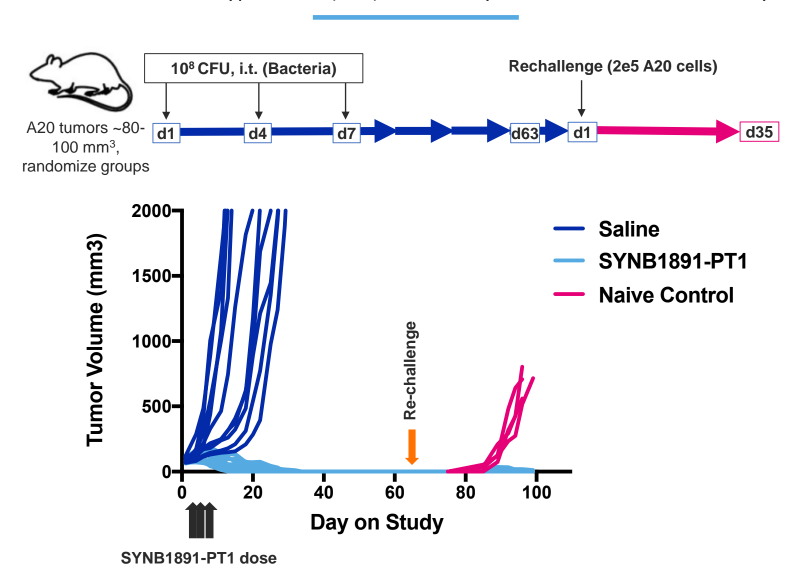
SYNB1891 In Vivo Characterization

Dose-dependent Anti-tumor Activity of SYNB1891 Prototype Strain (PT1) as a Single Agent



SYNB1891 In Vivo Characterization

SYNB1891 Prototype Strain (PT1) Leads to Systemic Anti-tumor Immunity





Dual Innate Immune Activator SYNB1891

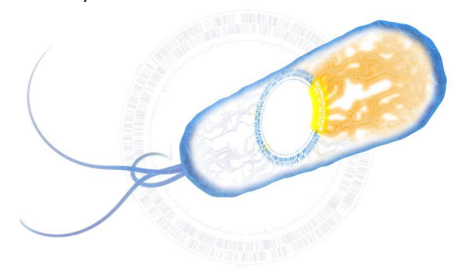
Designed to Locally Inflame the TME and Systemically Drive Tumor Antigen-Specific Immunity

PROGRESS TOWARDS THE CLINIC

- Tumor Colonization without Leakage
- Enhanced Activity vs. Naked STING Agonist
- Intracellular Activation of STING and Bacterial-Induced Immune Pathways Within APCs
- Dose-dependent Anti-tumor Activity
- Immunological Memory
- Clinical trial material manufactured
- IND Submission 2H19

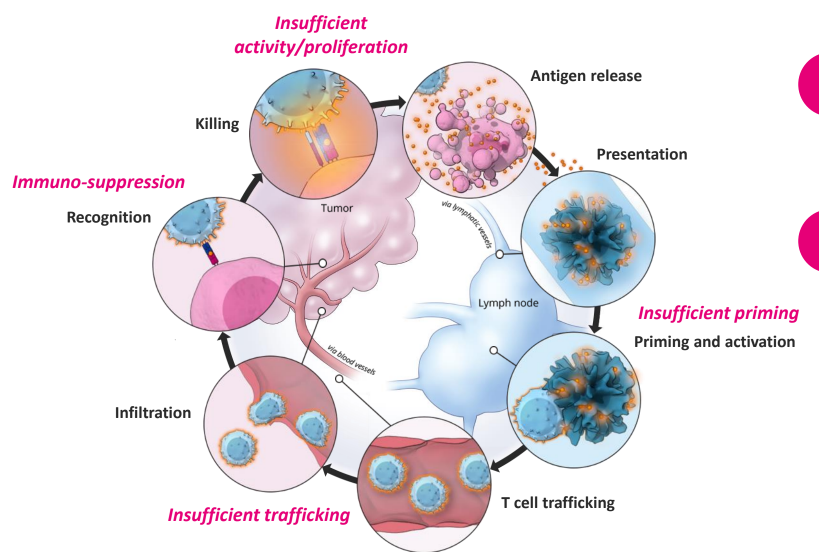
PROMISE OVER OTHER APPROACHES

- STING Agonism in Target Cells that Drive Efficacy
- Sparing Cells Where STING Agonism is Detrimental
- Activation of Multiple Innate Immune Pathways
- Low Systemic Risk





A Tumor Can Evade Multiple Critical Aspects of the Cancer-Immunity Cycle



MONOTHERAPIES OFTEN FAIL TO OVERCOME TUMOR EVASION MECHANISMS

Recognized Need to Combine Mechanisms to Broaden the Benefit of Immunotherapy

ENGINEER LIVING SOLUTIONS: SYNTHETIC BIOTIC MEDICINES

Rationally Designed for Combinatorial Effect

Locally Inflame the tumor microenvironment (TME)

Systemically Drive Tumor-Antigen Specific Immunity

In Situ Vaccination: Neo-antigen Priming and Sustained Immune Response

Additional Synthetic Biotic Effectors

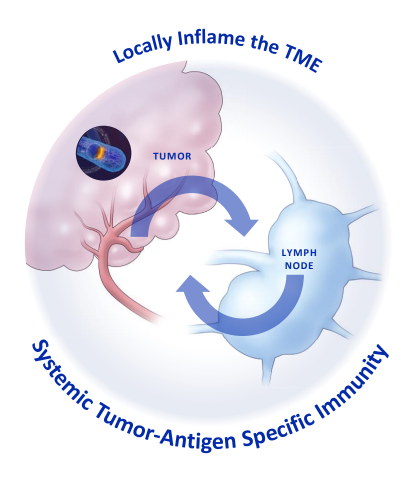
VISION: Rational Design to Locally Inflame the TME AND Systemically Drive Tumor-Antigen Specific Immunity

RELIEVE IMMUNOSUPRESSION

- Kyn Consumption
- Ade Consumption
- αPD-1 scFv

PROMOTE TRAFFICKING

- Chassis effect
- CXCL10
- Hyaluronidase



PROMOTE AND SUSTAIN IMMUNE ACTIVATION

- IL-15; IL-12
- Arg Production
- 4-1BBL
- OX40L

PRIME FOR TUMOR-ANTIGEN-SPECIFIC VACCINATION

- Chassis effect
- TNFα

• 5FC→5FU

IFNy

STING

- αCD47 ScFv / Sirpα
- αCD40 scFv/CD40L
 GM-CSF



Broad Ambitions in Immuno-Oncology

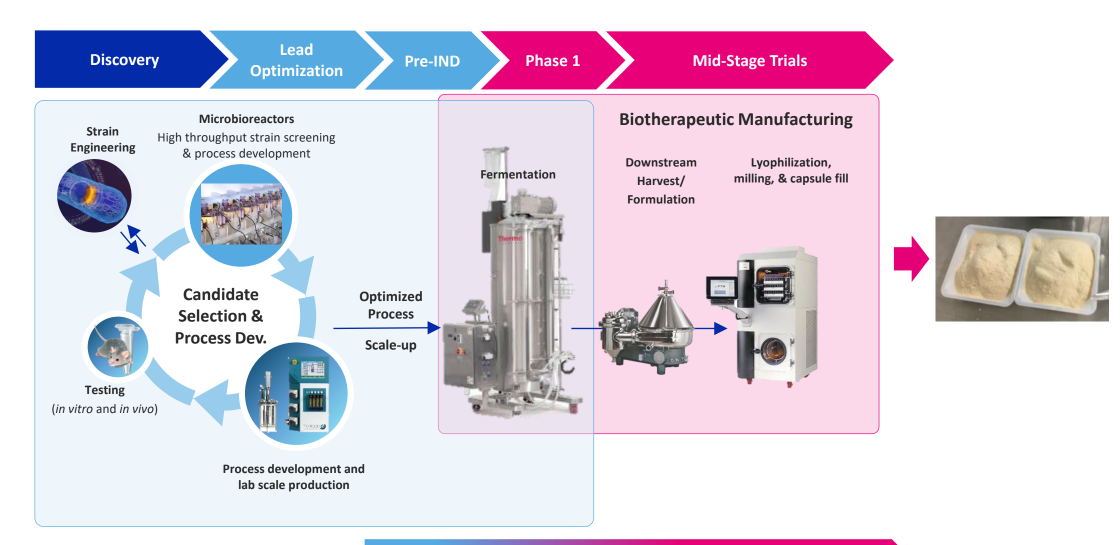
Vision: Expand and Exceed the Effect of Cancer Immunotherapies





Synlogic Internal GMP Manufacturing Capabilities

In-house Process Development and Clinical Manufacturing for Early & Mid-Stage Trials



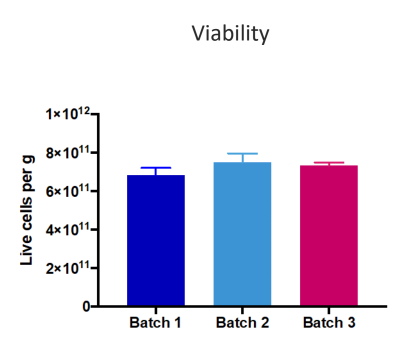
Analytical Methods Development and Validation

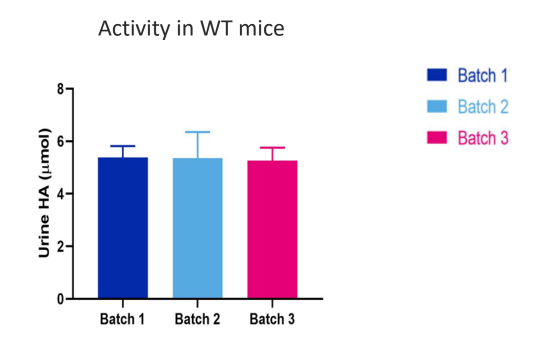
Demonstrated Progress in Development of Lyophilized SYNB1618

- Improved fermentation process enables production of a solid formulation of SYNB1618 with minimal impact on cell viability or activity
- Lyophilized SYNB1618 is similarly active to frozen liquid in terms of consumption of Phe or production of TCA/HA in vitro and in vivo
- New solid process material is expected to have improved quality attributes including less free protein and reduced viscosity
- Process is robust and reproducible at 30 L production scale
- Lyophilized SYNB1618 is stable for >90 days at 2-8 °C and >30 days at room temperature
- Suite build-out complete and ready to manufacture cGMP lyophilized SYNB1618



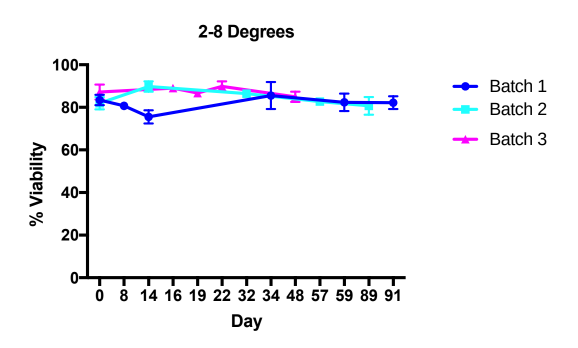
Batch to Batch Consistency of SYNB1618 Solid Formulation

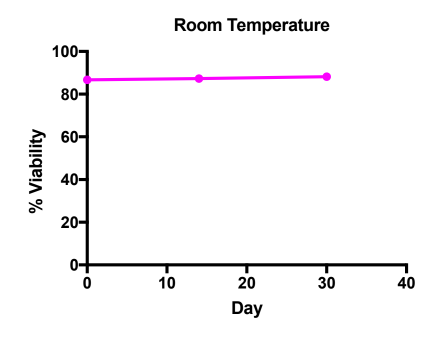






Stability of SYNB1618 Solid Formulation







Platform Collaboration to Accelerate Development of Synlogic's Synthetic Biotic Medicines



- Provides access to Ginkgo's industrial scale, highthroughput strain optimization and screening
- Enables screening and identification of higher quality optimized candidates, increasing potential for success



- Delivers novel tools for increased candidate potency
- Includes equity investment at a premium, extending runway through multiple milestones

Builds off validated pilot program initiated in 2017

Platform Collaboration to Accelerate Development of Synlogic's Synthetic Biotic Medicines



- Industry leader in the construction and editing of microbial strains and organisms
- Leaders in non-therapeutic commercial applications of synthetic biology
- Comprehensive database of microbial genome sequences and unparalleled automated foundry

Rapid prototyping and screening enables efficient iteration through 1000's of microbial strains

Top-tier platform companies and collaborations











High-quality investor base













2019 Progress and Milestones

SYNB1618 in PKU

- ✓ Completed Phase 1/2a study in healthy volunteers and patients, top-line data presented
 - Full data presentation Sept. 2019 (SSIEM) (safety, tolerability and biomarkers)

SYNB1020 in Hyperammonemia

- ✓ Preclin. and HV clin. data published in Sci. Transl. Med.
 - > Complete ongoing study in patients with cirrhosis
- Data expected 3Q2019 (safety, tolerability and ammonialowering)
- With ammonia-lowering data define development plan

SYNB1891 in Immuno-Oncology

- > IND submission 2H2019
- ✓ Clinical trial material manufactured
- ✓ Advance AbbVie collaboration expand Ginkgo collaboration
 - > Advance preclinical pipeline

