

Disclaimers

Forward Looking Statements

This communication contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. All statements contained in this communication that do not relate to matters of historical fact should be considered forward-looking statements, including statements about the potential benefits of any of our products or product candidates; the ultimate safety and efficacy data of SER-155; study results; plans to seek FDA feedback; clinical data and clinical trials; our intentions related to the development of SER-155; our intention to seek Breakthrough Therapy Designation; the ability of live biotherapeutics to prevent or reduce infections; or the timing of any of the foregoing and other statements which are not historical fact.

These forward-looking statements are based on management's current expectations. These statements are neither promises nor guarantees, but involve known and unknown risks, uncertainties and other important factors that may cause our actual results, performance or achievements to be materially different from any future results, performance or achievements expressed or implied by the forward-looking statements, including, but not limited to, the following: (1) we have incurred significant losses, are not currently profitable and may never become profitable; (2) our need for additional funding; (3) our history of operating losses; (4) the restrictions in our debt agreement; (5) our novel approach to therapeutic intervention; (6) our reliance on third parties to conduct our clinical trials and manufacture our product candidates; (7) the competition we will face; (8) risks associated with our clinical trials; (9) whether the FDA grants Breakthrough Therapy Designation; (10) our ability to protect our intellectual property; (11) our ability to eratin key personnel and to manage our growth; and (12) risks related to the proposed transaction under the Purchase Agreement with Société des Produits Nestlé S.A for the sale of the VOWST business to SPN. These and other important factors discussed under the caption "Risk Factors" in our Quarterly Report on Form 10-Q filed with the SEC, on August 13, 2024, and our other reports filed with the SEC could cause actual results to differ materially from those indicated by the forward-looking statements made in this communication. Any such forward-looking statements represent management's estimates as of the date of this communication. While we may elect to update such forward-looking statements at some point in the future, we disclaim any obligation to do so, even if subsequent events cause our views to change. These forward-looking statements should not be relied upon as representing our views as of any date subsequent to the date of this communication.



Disclaimers

Important Information About the Transaction and Where to Find it

In connection with the proposed transaction involving Seres Therapeutics, Inc. ("Seres") and Société des Produits Nestlé S.A. ("SPN"), Seres filed a definitive proxy statement with the Securities and Exchange Commission (the "SEC"). Seres may also file other relevant material with the SEC regarding the proposed transaction. Beginning on August 26, 2024, Seres mailed the definitive proxy statement to its stockholders. INVESTORS AND STOCKHOLDERS OF SERES ARE URGED TO READ THE DEFINITIVE PROXY STATEMENT AND OTHER RELEVANT MATERIALS CAREFULLY AND IN THEIR ENTIRETY WHEN THEY BECOME AVAILABLE BECAUSE THEY CONTAIN OR WILL CONTAIN IMPORTANT INFORMATION ABOUT SERES AND THE PROPOSED TRANSACTION. Investors may obtain a free copy of these materials (when they are available) and other documents filed by Seres with the SEC at the SEC's website at www.sec.gov or from Seres at its website at ir.serestherapeutics.com.

Participants in the Solicitation

Seres and certain of its directors, executive officers and other members of management and employees may be deemed to be participants in soliciting proxies from its stockholders in connection with the proposed transaction. Information regarding the persons who may, under the rules of the SEC, be considered to be participants in the solicitation of Seres' stockholders in connection with the proposed transaction is set forth in Seres' definitive proxy statement for its stockholder meeting, which was filed with the SEC on August 26, 2024, at which the proposed transaction will be submitted for approval by Seres' stockholders. You may also find additional information about Seres' directors and executive officers in Seres' Annual Report on Form 10-K for the fiscal year ended December 31, 2023, which was filed with the SEC on March 5, 2024, Seres' Definitive Proxy Statement for its 2024 annual meeting of stockholders, which was filed with the SEC on March 5, 2024, and in subsequently filed Current Reports on Form 8-K and Quarterly Reports on Form 10-Q.

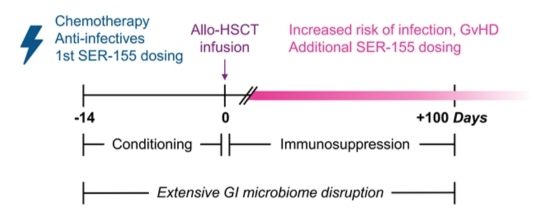


SER-155 is designed to reduce life-threatening complications of allo-HSCT



- Investigational live oral biotherapeutic cultivated from clonal master cell banks
- Designed to prevent GI-derived bacterial bloodstream infections (BSIs) and other pathogen-associated complications

Allo-HSCT treatment regimen and SER-155



- Only ~60% survival 3 years posttransplant
- ~10% transplant mortality in first
 100 days post-transplant
- ~80% of adult deaths in first 100 days caused by complications of procedure; half of these due to infections and GvHD
- Complications have substantial impact: mortality, cost, hospital stay



SER-155 Phase 1b study evaluated safety, pharmacology, and efficacy in adult allo-HSCT recipients

COHORT 1

Open-label (n=15 enrolled)

COHORT 2

Placebo-controlled 1:1 (n=45 enrolled)

SER-155

SER-155

Placebo

results reported May 2023

results announced Sept. 2024

Primary Endpoints:

- · Safety and tolerability
- SER-155 bacterial strain engraftment

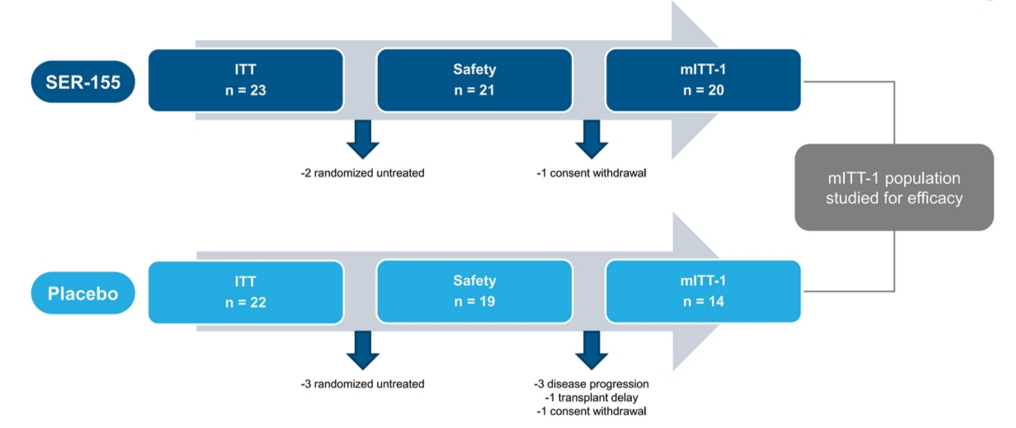
Key Secondary Endpoints through HSCT Day 100:

- Incidence of bloodstream infections (BSI), GI infections, and acute GvHD ≥ Grade 2
- · Incidence and duration of febrile neutropenia
- Bacterial pathogen abundance

Received US FDA Fast Track Designation in December 2023; Intend to pursue Breakthrough Therapy designation

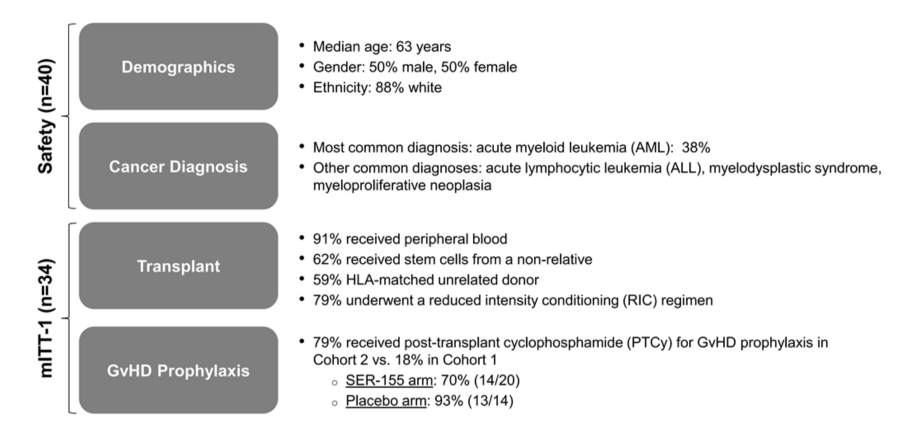


Cohort 2 patient disposition





Cohort 2 demographics: Representative of the allo-HSCT population





Patient Safety: Cohort 2 SER-155 was generally well tolerated with no treatment-related SAEs

Treatment-emergent adverse events (TEAEs)

- All but one subject in the placebo arm experienced at least 1 TEAE
- Most common for SER-155 treated subjects (≥50% and with Δ≥5% greater than placebo): diarrhea (86% vs. 74% placebo), nausea (62% vs. 53% placebo)
- 1/40 (3%) subject experienced a TEAE leading to treatment discontinuation (active = 0; placebo = 1)
- 3/40 (8%) subjects experienced a TEAE leading to study discontinuation (active = 1; placebo = 2)

Serious adverse events (SAEs)

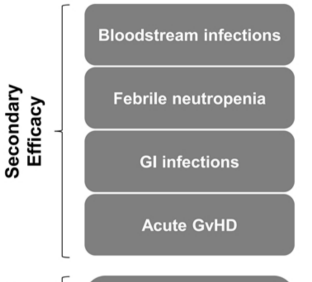
- 19/40 (48%) subjects experienced an SAE: 11/21 (52%) SER-155-treated subjects vs. 8/19 (42%) placebo-treated subjects; none considered related to SER-155 (no SUSARs)
 - Most common SAE SOC: infections & infestations (24% active vs. 37% placebo)
 - 3 deaths prior to Day 100 (active = 1; placebo = 2), 1 death after Day 100 (active), none considered related to SER-155

Adverse events of special interest (AESIs)

- AESIs (bloodstream infections, GI infection, invasive infection): 14/40 (35%) subjects
- Rates of AESIs were lower in SER-155 arm vs placebo arm (29% vs 42% respectively)
- No SER-155 species were identified in culture from any subject



Efficacy: SER-155 administration favorable with significant* reduction in both bacterial BSIs and systemic antibiotic exposure; lower febrile neutropenia



Significant decrease in bacterial bloodstream infections in SER-155-treated subjects vs. placebo

Numerically lower incidence rate of febrile neutropenia in SER-155-treated subjects vs. placebo

All GI infections were CDI**; 4 subjects in SER-155-treated (20%) and 2 subjects in placebo (14.3%) developed GI infections from HSCT Day 0-100

No subjects in either arm developed ≥ Grade 3 acute GvHD; 2 subjects in each arm developed Grade 2 acute GvHD

Exploratory

Exploratory

Efficacy

antimycotic exposure

Significantly lower mean cumulative exposure (days) to systemic antibacterials / antimycotics for SER-155-treated subjects vs. placebo

Significantly lower cumulative exposure rate to systemic antibacterials / antimycotics for SER-155-treated subjects vs. placebo

** CDI: C. difficile infection



^{*} no multiplicity adjustments were applied

Bloodstream infections from HSCT Day 0 to Day 100: Lower incidence in SER-155 treated subjects vs. placebo

Bloodstream infections from Day 0 to Day 100 (# patients)	SER-155 n=20 n (%)	Placebo n=14 n (%)
Subjects with confirmed BSI	2 (10.0%)	6 (42.9%)
95% confidence interval	(1.2, 31.7)	(17.7, 71.1)

mITT-1 population

Odds ratio	0.15
95% confidence interval	(0.01, 1.13)
p-value	0.0423

Organisms in SER-155 patients: Finegoldia magna; E. coli/Strep mitis

Organisms in placebo patients: E.coli; Enterococcus faecium/staph haemolyticus/Candida krusei; Staph aureus; Staph haemolyticus; Pseudomonas aeruginosa; E coli

- CI: 95% 2-sided Clopper-Pearson confidence interval of incidence is applied.
- Odds ratio: for incidence between treatment groups (SER-155 and placebo) with 95% 2-sided confidence interval and the corresponding p-value calculated based on the Fisher's Exact test



Febrile neutropenia from HSCT Day 0 to Day 100: Lower incidence in SER-155 treated subjects vs. placebo

Febrile neutropenia from Day 0 to Day 100 (# patients)	SER-155 n=20 n (%)	Placebo n=14 n (%)
Subjects with FN	13 (65.0)	11 (78.6)
95% confidence interval	(40.8, 84.6)	(49.2, 95.3)

mITT-1 population

Odds ratio	0.51
95% confidence interval	(0.07, 2.99)
p-value	0.4674



[.] CI: 95% 2-sided Clopper-Pearson confidence interval of incidence is applied

Odds ratio: for the incidence between treatment groups (SER-155 vs. placebo) with 95% 2-sided confidence interval and the corresponding p-value are calculated based on the Fisher's Exact test

Cumulative exposure to systemic antibacterials / antimycotics through HSCT Day 100:

Lower incidence in SER-155 treated subjects vs. placebo

Cumulative Antibacterial or Antimycotic Exposure (HSCT Days)	SER-155 n=20 n (%)	Placebo n=14 n (%)
Mean (SD)	9.2 (5.44)	21.1 (20.31)
Median	9.0	14.0
Min, Max	0, 19	0, 74

Mean Difference (95% CI)	-11.9 (-23.85, -0.04)
p-value	0.0494

mITT-1 population

 ^{95%} confidence interval and p-value based on independent samples t-test of the difference in mean days between SER-155 and placebo



Cumulative exposure is the sum of all days a subject received systemic antibacterials and/or antimycotics between HSCT Day 0 through Day 100; counting once per day regardless of number of agents taken

Cumulative exposure rate to systemic antibacterials / antimycotics through HSCT Day 100:

Lower incidence in SER-155 treated subjects vs. placebo

Cumulative Antibacterial or Antimycotic Exposure Rate (% of days)	SER-155 n=20 n (%)	Placebo n=14 n (%)
Mean (SD)	0.090 (0.0530)	0.305 (0.2898)
Median	0.089	0.244
Min, Max	0.00, 0.18	0.00, 0.90
		mITT-1 population

Mean Difference (95% CI) -0.2 (-0.38, -0.05)
p-value 0.0163

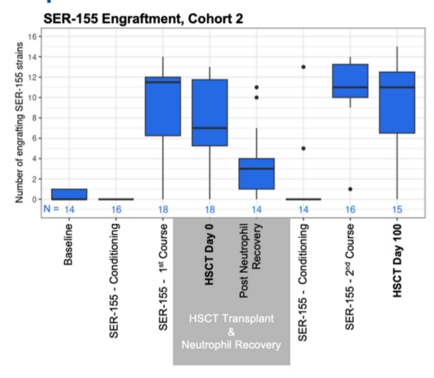
mITT-1 population



Cumulative exposure rate is calculated as the sum of all days a subject received systemic antibacterials and/or antimycotics on or after HSCT Day 0 (counting once per day, regardless of number of antibacterial/antimycotic medications taken in a day) through HSCT Day 100 over the total number of days a subject was on the study from HSCT Day 0 to the earliest of EOS, or HSCT Day 100

 ^{95%} confidence interval and p-value are based on independent samples t-test of the difference in mean days or mean rate of cumulative exposure between SER-155 and Placebo

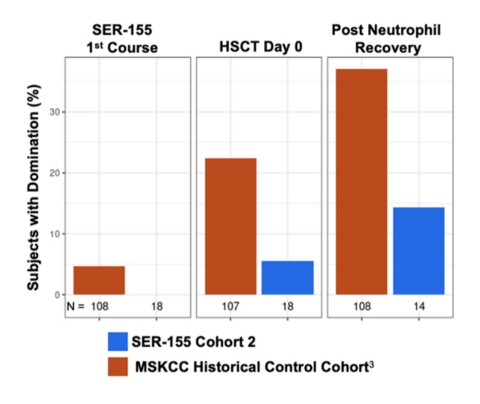
SER-155 Strain Engraftment: Primary objective achieved - drug bacteria strain engraftment was robust and as expected



- The majority of SER-155 strains were present at start of HSCT conditioning and durable through chemotherapy exposure
- Engraftment decreased but was detectable postneutrophil recovery, suggesting sustained engraftment, even under unfavorable GI conditions (e.g., antibiotic exposure), and through period of greatest BSI susceptibility
- The second course of SER-155 was effective at increasing strain engraftment following transplant & neutrophil recovery, with engraftment durable out to day 100 following transplant
- Cohort 1 and Cohort 2 engraftment magnitude and kinetics had high congruence



Pathogen Domination: Prevalence in SER-155 Cohort 2 was substantially lower relative to Historical Control Cohort



- SER-155 was designed to reduce pathogen domination that has been associated with risk of BSIs and other negative clinical outcomes¹
- In Cohort 2, the ability to detect pathogen domination² (i.e., relative abundance in the GI ≥30%) in the placebo arm, and differences between the study arms, was constrained due to the limited number of placebo stool samples and an imbalance in the number of available stool samples between the arms
- Observed pathogen domination events were low in the placebo and SER-155 arms with no significant differences observed
- Pathogen domination was substantially lower in SER-155 Cohort 2 compared to Historical Control Cohort³



Summary and Next Steps

Seres to engage with FDA on advancement of SER-155 allo-HSCT program

- Seek Breakthrough Therapy designation given high unmet medical need associated with bloodstream infections
- Pursue additional designations (Orphan Drug designation, Qualified Infectious Disease Product)

Phase 1 results support Seres' strategy to pursue SER-155 and other live biotherapeutics for prevention of serious bacterial infections

- Intend to evaluate SER-155 in additional patient populations with high risk of serious bacterial infections
- Opportunities to address multiple medically vulnerable patient groups with SER-155 and additional pipeline programs

