

Trial in progress: A phase 1 trial of TSC-100 and TSC-101, engineered T cell therapies that target minor histocompatibility antigens to eliminate residual disease after hematopoietic cell transplantation



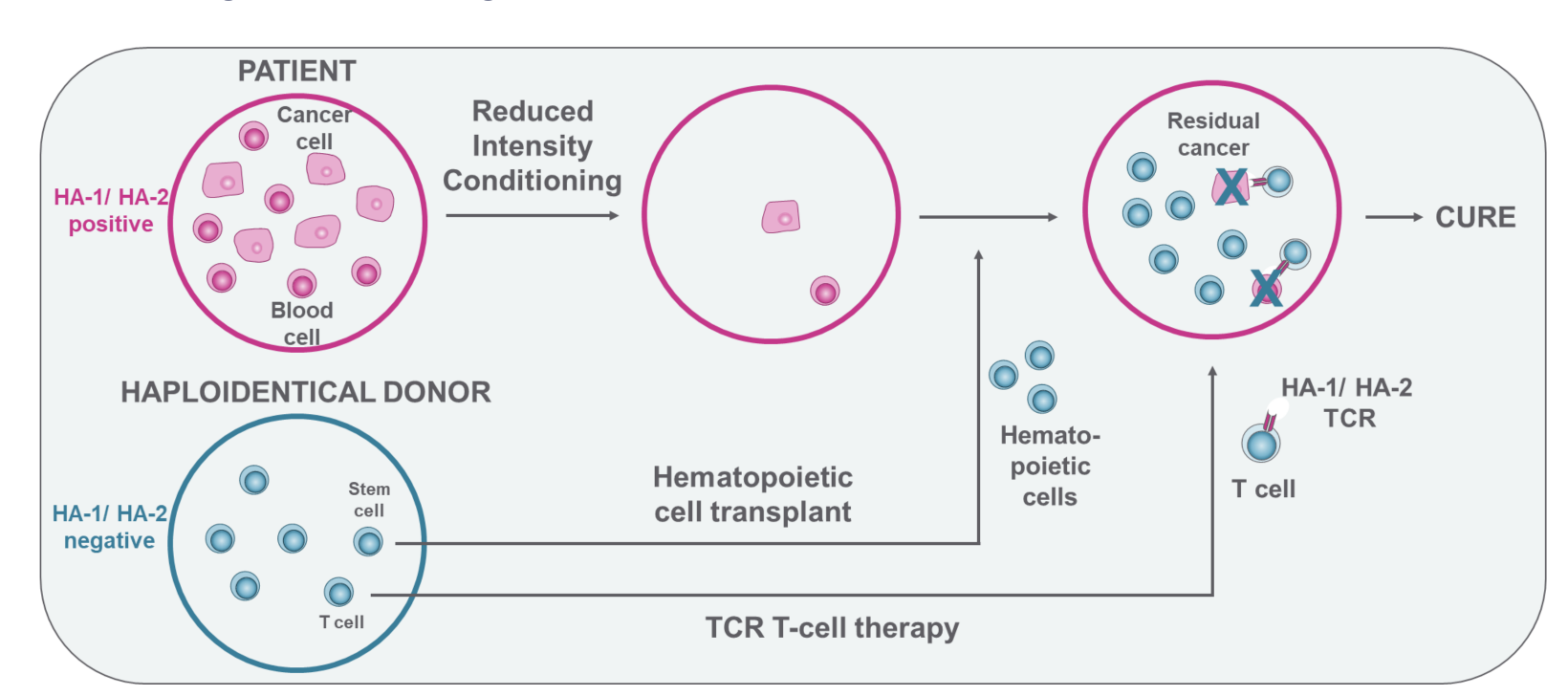
Monzr Al Malki<sup>1</sup>, Alla Keyzner<sup>2</sup>, Hyung C. Suh<sup>3</sup>, Uday Popat<sup>4</sup>, Saar Gill<sup>5</sup>, Yi-Bin Chen<sup>6</sup>, Melhem Solh<sup>7</sup>, Joseph Uberti<sup>8</sup>, Lohith Gowda<sup>9</sup>, Erica Buonomo<sup>10</sup>, Yun Wang<sup>10</sup>, Jim Murray<sup>10</sup>, Gavin MacBeath<sup>10</sup>, Debora Barton<sup>10</sup>, Shrikanta Chattopadhyay<sup>10</sup>, Ran Reshef<sup>11</sup>

Abstract # CT151

<sup>1</sup>City of Hope Medical Center, Duarte CA; <sup>2</sup>Mount Sinai Hospital, New York NY; <sup>3</sup>Hackensack University of Pennsylvania, Philadelphia PA; <sup>6</sup>Massachusetts General Hospital, Boston MA; <sup>7</sup>Northside Hospital, Atlanta GA; <sup>8</sup>Karmanos Cancer Institute, Detroit MI; <sup>9</sup>Yale University, New Haven CT; <sup>10</sup>TScan Therapeutics, Waltham MA; <sup>11</sup>Columbia University, New York NY

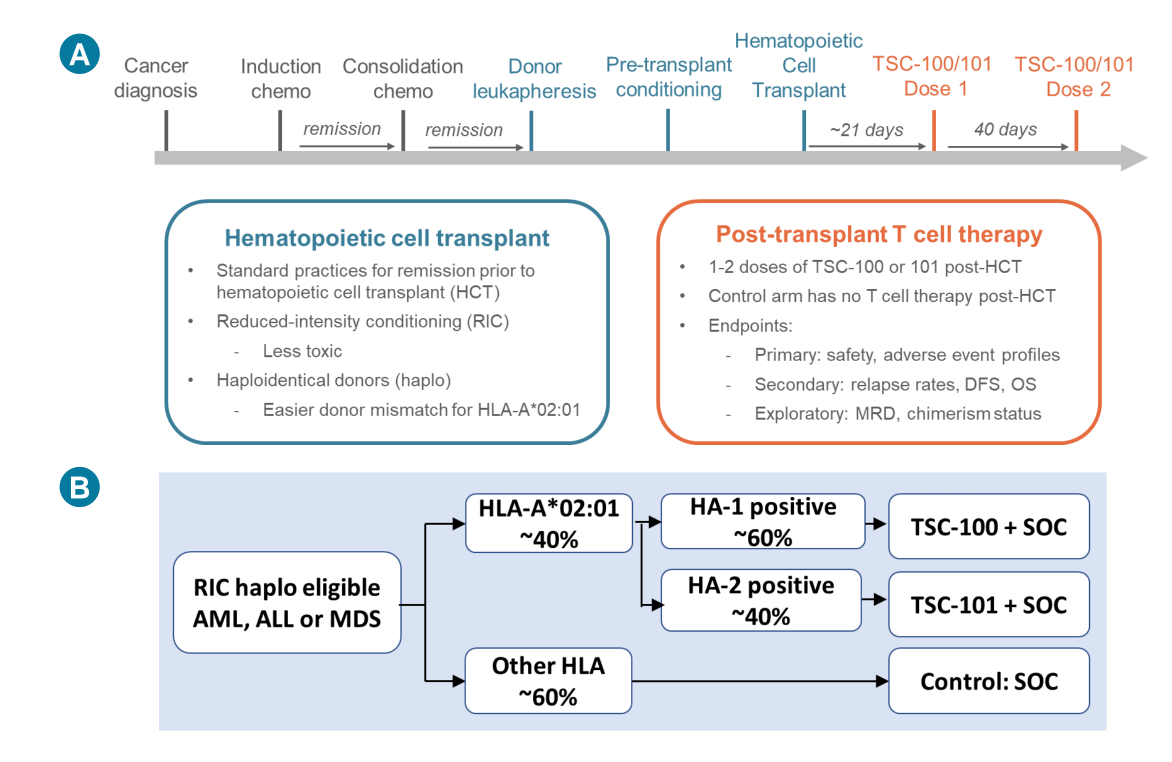
# **Background and rationale**

- Engineered T cell therapies have been transformative for lymphoid malignancies because depleting normal B cells or plasma cells can be tolerated. Other hematologic malignancies have not benefited because depleting other normal blood cells like myeloid cells cannot be tolerated.
- Allogeneic hematopoietic cell transplantation (HCT) remains the best curative option for most hematologic malignancies, yet relapses occur in ~40% of patients post-HCT and are associated with high mortality.
- A potential solution is targeting hematopoietic-lineage specific minor histocompatibility antigens (MiHAs) mismatched between transplant recipients and their donors.
- TScan has developed the engineered T cell products TSC-100 and TSC-101 that express TCRs targeting MiHAs HA-1 and HA-2 respectively, both presented by HLA-A\*02:01 and expressed only in hematologic cells.
- By choosing HCT patients who are HLA-A\*02:01 positive and either HA-1 or HA-2 positive, and donors who are mismatched on either MiHA or HLA-A\*02:01, TSC-100 and TSC-101 can eliminate all residual recipient hematologic cells while leaving donor hematologic cells untouched.



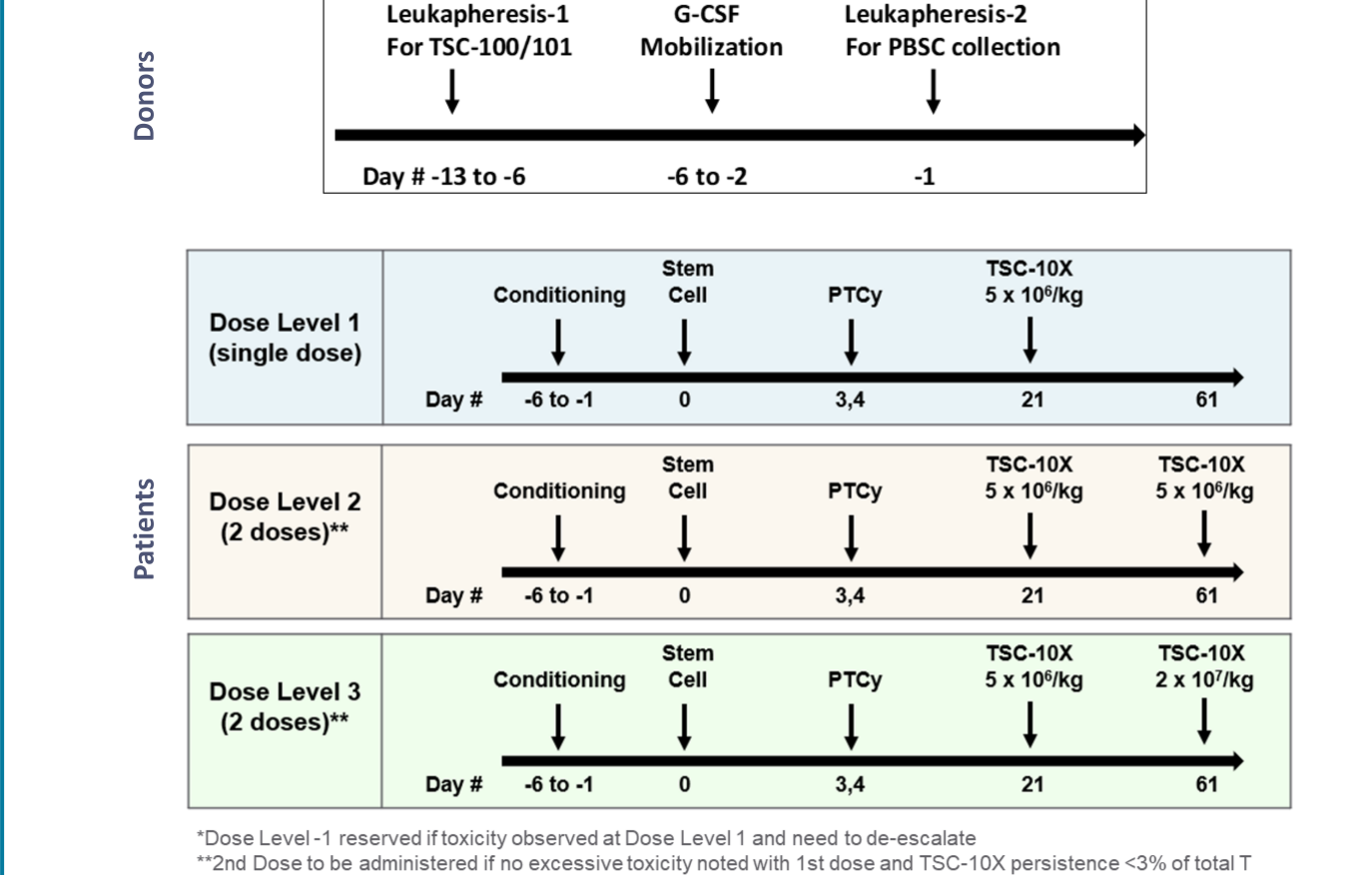
#### TSC-100 and TSC-101 target HA-1 and HA-2 **HA-1 TCR** Promoter 200 pg/mL 66 pg/mL **Promoter** 0.2 pg/mL **Modifications to** Achieves strong contribution of CD4 of engineered T and stable constant region T cells in clinical cells and tracking in expression of our ensure surface engineered TCR expression and product, boosting the patient **HA-2 TCR** proper TCR pairing CD8<sup>+</sup> T cell function - 10 **p**g/mL NTC <u>→</u> 1 ng/mL 100 pg/mL **Left**: Common vector used to manufacture both TSC-100 and TSC-101. **Right**: potencies of TSC-100 (A) and TSC-101 No peptide (B) were measured using peptide-pulsed T2 cells vs. non-targeted control TCRs. Time (h)

# Treatment plans and assignment to treatment arms



(A) All participants receive standard of care (SOC, blue) and treatment arm participants receive investigational (orange) treatment for patients with AML, ALL or MDS undergoing hematopoietic cell transplantation (HCT) following reduced intensity conditioning (RIC) from a haploidentical donor (haplo). (B) RIC-haplo eligible patients are assigned to treatment or control arms depending on their HLA and HA-1/ HA-2 genotypes and receive TSC-100/101+ SOC or SOC alone.

# Investigational treatment plans and dose escalation cohorts



Investigational treatment plans for donors (top) or patients (bottom). Donors undergo two rounds of leukapheresis, first before G-CSF mobilization, to manufacture TSC-100/101, and second after mobilization, for standard peripheral blood stem cell collection. Patients receive conditioning therapy from Days -6 to -1, stem cell infusions on Day 0, post-transplant cyclophosphamide (PTCy) on Days 3,4 then upon count recovery (around Day 21), receive the 1st dose of TSC-100 or 101. In Dose Levels 2 and 3, a second dose is administered at least 40 days after the 1st dose. In Dose Level 3, the 2nd dose is escalated by 4-fold. Dose escalation rules follow the interval 3+3 design1 with 1-12 patients per cohort. The study is currently enrolling patients at Dose Level 3.

cells, after review by the SRC and notification of FDA.

# Inclusion/ exclusion criteria and key specifications for study NCT05473910

**Exclusion Criteria** 

# **Inclusion Criteria**

# Patients in all arms:

- ≥18 years with AML, ALL or MDS
- ECOG-PS ≤2 any time in screening period • Eligible for reduced intensity conditioning
- Eligible for haploidentical donor HCT
- Treatment arms: HLA-A\*02:01 positive
- TSC-100 arm: HA-1+/- or HA-1+/+
- TSC-101 arm: HA-2+/- or HA-2+/+ Agree with 15-year long term follow up
- Control arm: Any HLA type apart from HLA-A\*02:01 or HLA-A\*02:01 positive without suitably mismatched donor

### **Donors in treatment arms:**

- ≥ 18 years old
- Able to undergo peripheral blood stem cell (PBSC) collection & 2 rounds of leukapheresis
- Donors matched to TSC-100 participants should be negative for all HLA-A\*02 alleles or HA-1-/- (negative)
- Donors matched to TSC-101 participants should be negative for all HLA-A\*02 alleles

 Levels of donor-specific HLA antibodies high enough to warrant desensitization protocols

and who have no alternate donors

- Treatment arms: HLA-A\*02:07 positive Patients with evidence of clinically
- significant infection or uncontrolled viral reactivation of cytomegalovirus (CMV), Epstein-Barr virus (EBV), Adenovirus, BK virus (BKV), or human herpesvirus 6 (HHV-6)
- Prior allogeneic HCT

# **Donors in treatment arms:**

Patients in all arms:

- Donors for TSC-100 positive for any HLA-A\*02 allele, unless they are HA-1 negative. Donors for TSC-101 positive for any HLA-
- A\*02 allele regardless of HA-2 status. Donors who test positive for: HIV-1, HIV-2, HTLV-1, HTLV-2 or with active hepatitis B or
- hepatitis C, syphilis, West Nile virus infectior or screen positive for risk of Creutzfeldt-Jakob disease or Zika virus with questionnaires.

**RIC regimens:** 

 Fludarabine/ cyclophosphamide/ total body irradiation (200 or 400 cGy)

**Protocol Specifications** 

- Fludarabine/ melphalan +/- / total body irradiation (200 cGy)
- Thiotepa/ busulfan/ fludarabine
- Fludarabine/ melphalan/ thiotepa

## **GvHD** prophylaxis:

- Post-transplant cyclophosphamide (Days
- Mycophenolate (until >Day 35)
- Tacrolimus (until >Day 90)

#### **Acute or chronic GvHD treatment:**

Any therapy per institutional guidelines

### **Maintenance therapies:**

- FLT3, BCR/Abl, IDH inhibitors- allowed 60
- days post TSC-10X or after Day 100 Other anti-leukemia agents (e.g. oral azacytidine) not allowed

# **Exploratory endpoints of minimal residual disease (MRD) and donor chimerism can** indicate biological activity and early efficacy

## **Minimal Residual Disease**

- Combination of next-generation sequencing (NGS) with flow cytometry detects MRD in 40% of AML patients<sup>2</sup>.
- Pre-transplant MRD+ patients have ~67% risk of relapse

- Post-HCT MRD+ by flow alone tends to be low ~16%4. NGS expected to double MRD detection<sup>2</sup>. Post-HCT MRD+ patients have up to 90% chance of

# MRD detection approach

- MRD will be detected in pre- and post-transplant bone marrow biopsies with a combination of flow (local sites) and NGS (central lab)
- Conversion from MRD positive to negative can be an early indicator of biological activity and an early surrogate of efficacy

# 

**HA-positive residual** 

malignant cells

# **HA-positive residual**

# Standard STR-based assay

# • Pros: clinically validated; measurable in all patients; mixed

chimerism predicts ~60% risk of relapse<sup>6</sup> Cons: Poor limit of detection (~1%); PTCy causes high donor

Mixed donor cell chimerism

chimerism > 98% by Day 30<sup>7</sup>

## Novel NGS-based assay (AlloHeme)

- Pros: NGS of ~400 SNPs improves limit of detection to
- Cons: Predictive value of NGS assay unknown, trial ongoing (NCT04635384)

## **Chimerism detection approach:**

- Chimerism will be detected in bone marrow, whole blood, CD3 and CD33 subsets using standard STR and novel NGS
- Complete donor chimerism and faster kinetics can indicate biological activity and early efficacy

**Endpoints**: Primary endpoints include adverse event profile and dose limiting toxicities. Secondary endpoints include relapse rates at 1 year, disease-free-survival and overall survival. Exploratory endpoints include complete or mixed donor chimerism rates and kinetics, MRD+ rates before and after HCT and TSC-100/101 persistence in the treatment arms. Following transplantation, residual HA-positive patient-derived malignant cells are measured with high-sensitivity MRD assays (left) whereas residual HA-positive patient-derived hematologic cells, malignant, pre-malignant or normal, are measured using standard and high-sensitivity chimerism assays (left). Clearance of MRD or mixed donor cell chimerism can be early indicators of biological activity and early surrogates of efficacy. Preliminary results were presented at ASH, Dec 20238 and the Best Abstracts session at the Tandem Transplantation and Cellular Therapy Meeting, Feb 2024 with abstract available at: https://tandem.confex.com/tandem/2024/meetingapp.cgi/Paper/23846

hematologic cells

# REFERENCES

- 1. Liu M et al. J Biopharm Stat. 2020 . Jongen-Lavrencic M et al. N Engl J Med. 2018
- 4. Craddock C et al. J Clin Oncol. 2021

7. Legrand F et al. Blood 2016 8. Al Malki M et al, Blood. 2023