FEBRUARY 2020

Next©ure

Next-Generation Immunomedicines

Forward-Looking Statements

To the extent that statements contained in this presentation are not descriptions of historical facts, they may be deemed to be forward-looking statements under the Private Securities Litigation Reform Act of 1995. Words such as "may," "will," "expect," "anticipate," "estimate," "intend," "next," "near-term," "future" and similar expressions, as well as other words and expressions referencing future events, conditions, or circumstances, are intended to identify forward-looking statements. Examples of forward-looking statements in this presentation may include, among others, statements regarding: (i) the timing, progress and results of our preclinical and clinical trials; (ii) the timing or likelihood of regulatory filings for our product candidates; (iii) our manufacturing capabilities and strategy; (iv) the potential benefits and activity of our product candidates; (v) our expectations regarding the nature of the biological pathways we are studying; (vi) our expectations regarding our FIND-IO platform; and (vii) the potential benefits of our relationships with Dr. Lieping Chen and Yale University.

Various factors could cause actual results to differ materially from those projected in any forward-looking statement. Such risks and uncertainties include, among others: our limited operating history and no products approved for commercial sale; our history of significant losses; our need to obtain additional financing; risks related to clinical development, marketing approval and commercialization; and the unproven approach to the discovery and development of product candidates based on our FIND-IO platform. No forward-looking statement is a guarantee of future results or events, and one should avoid placing undue reliance on such statements. For further discussion of these and other factors that could affect the outcome of our forward-looking statements, see our filings with the Securities and Exchange Commission, including in "Risk Factors" and "Special Note Regarding Forward-Looking Statements" in the Risk Factors section and throughout NextCure's Form 10-Q filed with the SEC on November 12, 2019. Except as otherwise indicated, this presentation speaks as of the date indicated herein. Except as required by law, we assume no obligation to update any forward-looking statements, or to update the reasons why actual results could differ materially from those anticipated in the forward-looking statements, even if new information becomes available in the future. The information in this presentation is not complete and may be changed.





Unmet Medical Needs of Cancer Patients



We Need New Solutions



Product Development Pipeline

PROGRAMS	CELLS	DISCOVERY	PRECLINICAL	PHASE 1	PHASE 2	PHASE 3	NEXT MILESTONE	WORLDWIDE RIGHTS
PRODUCT CANDIDATES								
NC318 (S15) Monotherapy	Tumors and macrophages	ONCOLOGY	/				Phase 2 data by end of Q4 2020	Next© ure
NC318 (S15) Chemo Combo	Tumors and macrophages	ONCOLOGY					Initiate Phase 1 mid-2020	Next© ure
NC410 (LAIR-1)	Dendritic and T cells	ONCOLOGY					IND filing in Q1 2020	Next© ure
DISCOVERY AND RESEARCH PROGRAMS								
Multiple Programs	lmmune cells						First IND filing in early 2021	Next© ure
FIND-IO Platform	Multiple cell types						First IND filing in late 2022	Next© ure

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NC318 Humanized Monoclonal Antibody



TARGET Siglec-15 ("S15")

CELL TYPES Tumors &

macrophages

MOA Blocks S15-induced immunosuppression

INDICATIONS

NSCLC, ovarian, head & neck and triple negative breast cancers

S15 as a Target

EXPRESSION

Tumors and Macrophages



medicine

Wang et al.,

2019

FUNCTION

Potently Suppresses T Cell Function



NON-RESPONDERS

Generally Non-Overlapping with PD-L1 Expression



Siglec-15 as an immune suppressor and potential target for normalization cancer immunotherapy

ARTICLES https://doi.org/10.1038/s41591-019-0374-x

NC318 Phase 1 Trial Status as of November 9, 2019 (SITC Presentation) DOSE ESCALATION AND SAFETY AND TOLERABILITY

Completed

ENROLLMENT

- 49 patients
- 15 tumor types
- Median of 3 prior therapies
- All comers regardless of PD-L1 or S15 expression status

SAFETY

- No DLTs through 800 mg
- 1 DLT at 1600 mg: Grade 3 pneumonitis
- Common irAEs observed, including diarrhea, rashes, vitiligo, arthralgias

Landone

RESPONSES

- Evaluations every 8 weeks
- 1 confirmed CR (55+ weeks)
- 1 confirmed PR (28+ weeks)
- 14 durable SD (≥16 weeks)







Yale University

Most common AEs: infusion reactions, fatigue, headaches, pruritis, elevated amylase and elevated lipase



NC318: Single-Agent Activity in PD-1 Refractory NSCLC

BASELINE WEEK 16 Confirmed Image: Complete response

Target lesion gone

56 y/o NSCLC with multiple lesions (PD-L1 TPS <50%) 8 mg every 2 weeks

PRIOR THERAPIES:

Chemotherapy (x3) Nivolumab (best response stable disease then progression)

Confirmed

PARTIAL RESPONSE

2.93 cm

Target lesion



Target lesions -71%

74 y/o NSCLC (PD-L1 TPS <50%) 400 mg every 2 weeks

LAST PRIOR THERAPY:

Immunotherapy: LAG3/PD-1 (best response stable disease then progression)



NC318 Phase 2 Trial Status as of November 9, 2019 DOSE EXPANSION - ENROLLING



NC410 Decoy Human Fusion Protein Targeting the TME



TARGET

Leukocyte-Associated Immunoglobulin-like Receptor-1 (LAIR-1) CELL TYPES Dendritic cells and T cells

MOA Promotes T cell function & dendritic cell activity

INDICATIONS

Advanced or metastatic solid tumors

NC410 Prevents Immune Suppression

NC410 IS A FUSION PROTEIN OF LAIR-2 AND A DECOY FOR LAIR-1 AND PROMOTES T CELL FUNCTION AND DC ACTIVATION







NC410 Summary

Promotes T cell function and dendritic cell activity in preclinical studies

IND-enabling tox studies complete



cGMP manufacturing complete

IND filing expected Q1 2020





Finding Solutions with a Powerful Discovery Engine

Functional, Integrated, NextCure Discovery in Immuno-Oncology





Anticipated Near-Term Milestones



