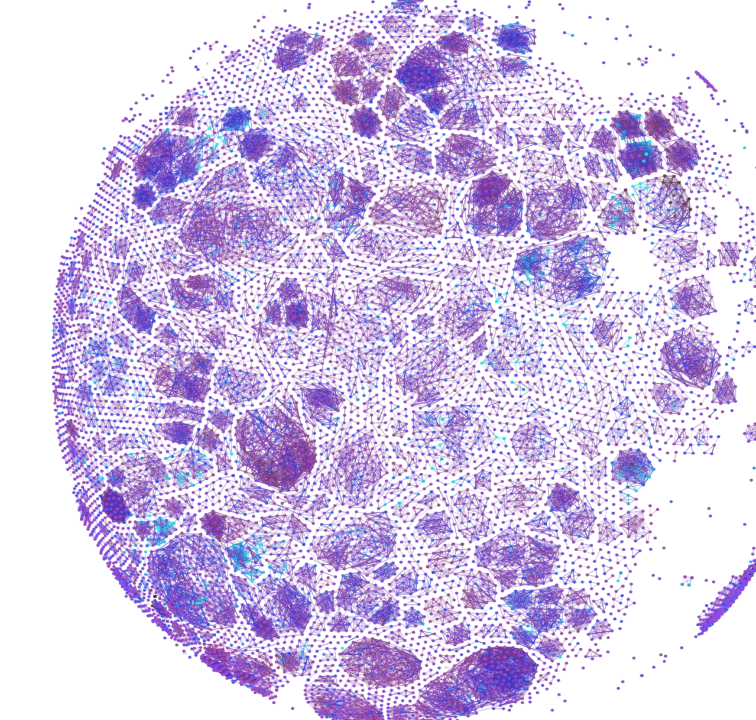
## Decoding Biology To Radically Improve Lives

End of Q4 2022







## **Disclaimers**

This presentation and any accompanying discussion and documents contain information that includes or is based upon "forward-looking statements" within the meaning of the Securities Litigation Reform Act of 1995. These forward-looking statements are based on our current expectations, estimates and projections about our industry and our company, management's beliefs and certain assumptions we have made. The words "plan," "anticipate," "believe," "continue," "extimate," "expect," "intend," "may," "will" and similar expressions are intended to identify forward-looking statements. Forward-looking statements made in this presentation include statements about Recursion's use of the proceeds from its private placement, the initiation, timing, progress, results, and cost of our research and development programs and our current and future preclinical and clinical studies, the potential size of the market opportunity for our drug candidates, our ability to identify viable new drug candidates for clinical development and the accelerating rate at which we expect to identify such candidates, our expectation that the assets that will drive the most value for us are those that we will identify in the future using our datasets and tools, and many others. Forward-looking statements made in this presentation-are neither historical facts nor assurances of future performance, are subject to significant risks and uncertainties, and may not occur as actual results could differ materially and adversely from those anticipated or implied in the forward-looking statements. For a discussion of factors that could affect our business, please refer to the "Risk Factors" sections in our filings with the U.S. Securities and Exchange Commission, including our Annual Report on Form 10-K. This presentation does not purport to contain all the information that may be required to make a full analysis of the subject matter. We undertake no obligation to correct or update any forward-looking statements, whether as a result of new information, future e

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## Maturing the TechBio value proposition in 2022

**Initiated 5 clinical trials** in 2022 (3 Ph2, 2 Ph1) and planning to initiate a **6**<sup>th</sup> **clinical trial** (Ph1b/2) for AXIN1 or APC mutated oncology in early 2024

Expecting REC-3964 Ph1 readout in 2H 2023, REC-994 Ph2 top-line data in 2H 2024, and REC-2282 Ph2 interim analysis in 2024

**Novel oncology programs** (RBM39, Target Alpha) nearing **IND-enabling studies** 

Advancing collaborations in **Fibrosis (Bayer)** and **Neuroscience (Roche-Genentech)** 

 \$13B in potential milestones across 50+ possible programs plus royalties

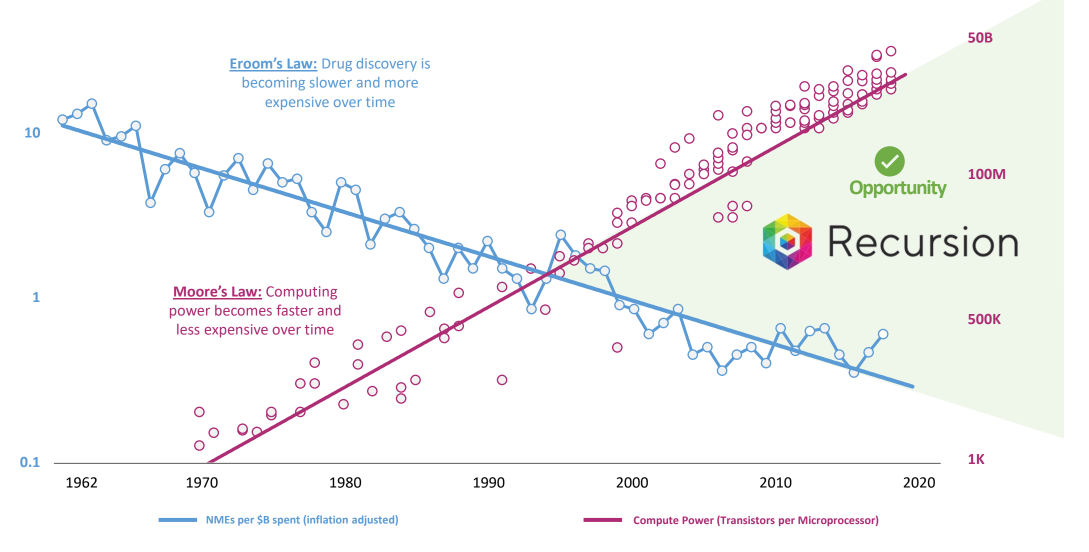
We believe that we have built one of the largest proprietary & relatable in-vitro biological and chemical datasets on Earth

>21 petabytes of data and>3 trillion searchable relationships





# Recursion has an opportunity for arbitrage at the intersection of technology and biology



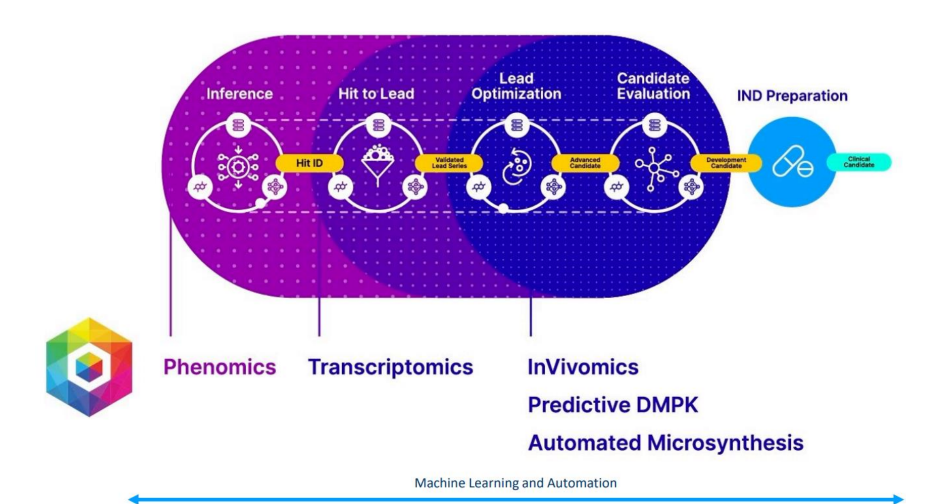


# Recursion's map-based approach is designed to set the standard for drug discovery in the 21st century

Traditional Drug Discovery		Recursion Approach		
	<b>Literature</b> drives discovery. <i>Informs target-based hypotheses</i>	VS	*	Platforms drive discovery.  Unbiased & target agnostic
	<b>Data</b> are an exhaust. <i>Limited to testing hypotheses</i>	VS		Data are our fuel. Shape our hypotheses
	<b>Disparate data</b> generation.  Siloed to individual programs and diseases	VS		Connected data across programs. Relatable high-dimensional data
$\Leftrightarrow$	<b>Linear process</b> .  Little cross-program learning or iteration	VS		Virtuous cycles of atoms & bits. Iterative feedback accelerates learning
0 0	Bespoke processes.  Low-dimensional assays & biomarkers	VS	<u>≅</u> *	Industrialized to scale.  Automation & standardization

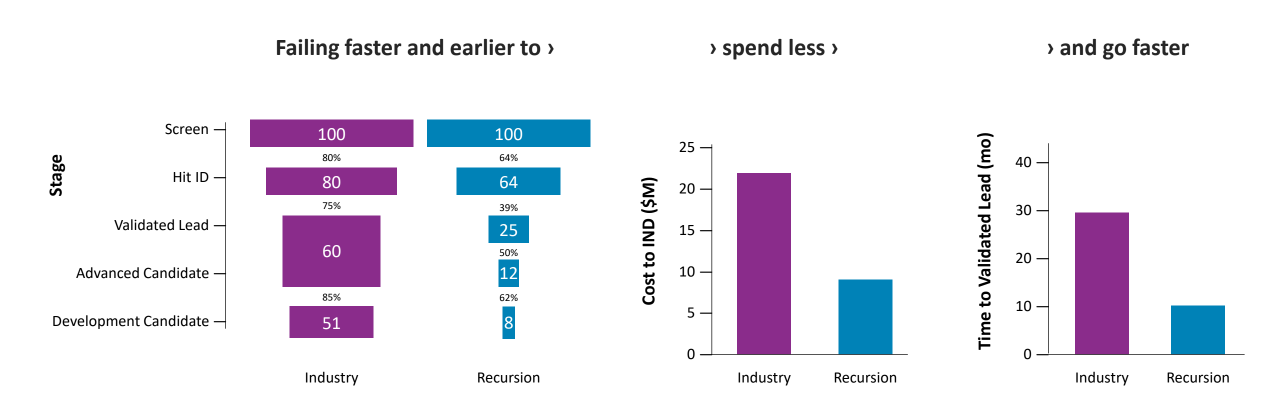


## How we aim to industrialize drug discovery



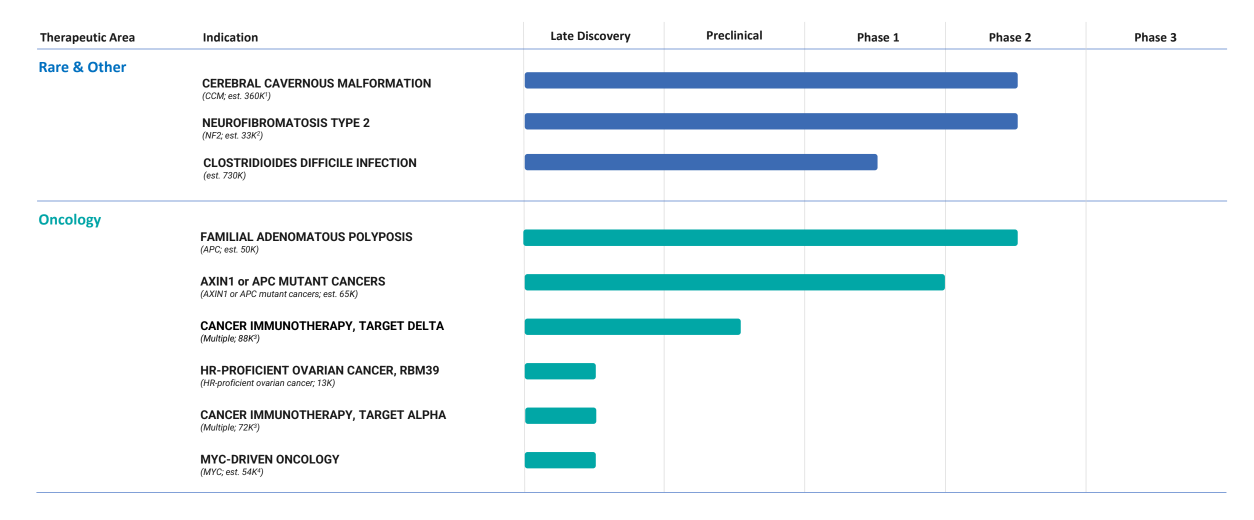


## Mapping and navigating the complex systems of biology and chemistry has demonstrated leading indicators of efficiency



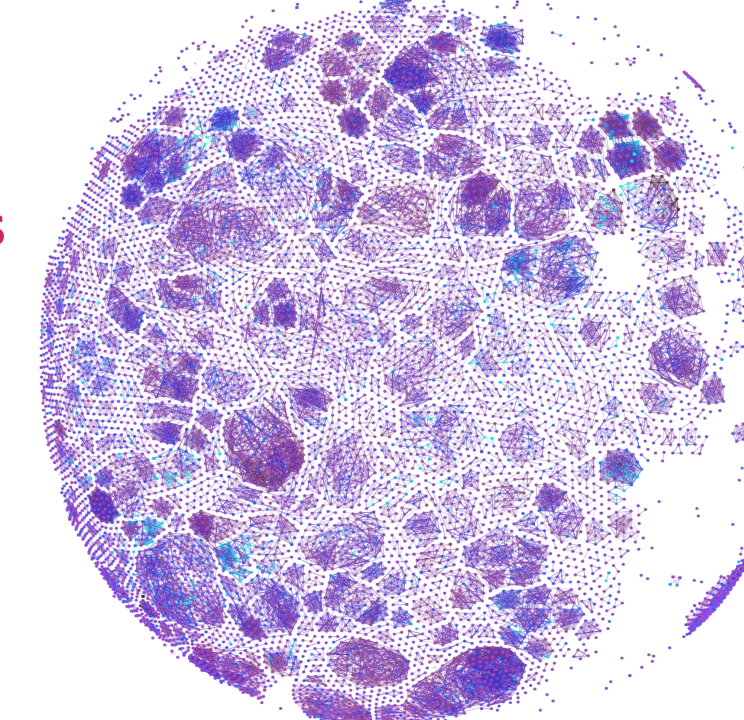


## Our pipeline reflects the scale and breadth of our approach



More than a dozen early discovery and research programs in oncology, neuroscience, inflammation & immunology, and rare disease

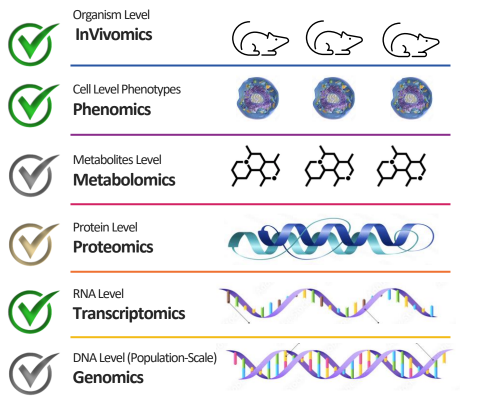
How we build maps of biology and chemistry to turn drug discovery into a search problem







## We build biological and chemical datasets to map relationships across scales and understand the connectivity of the system











Like digital maps of Earth, connections within and between layers add useful context. Similarly, Recursion is mapping different multiomic layers of biology and identifying connections within and between layers to better understand biology at scale.



#### **Robotic Automation at Scale**

Up to 2.2 Million wet-lab experiments per week profiling genes and compounds, we believe we are one of the largest phenomics (human cellular image-based) data producers





### **Digitization of Biology and** Chemistry

>21 Petabytes of proprietary highdimensional data, we believe this is one of the largest relatable in vitro biological and chemical datasets

#### **Diverse Biological and Chemical Inputs**

48

different human cell types

~1.7 Million

small molecule library, we believe this scale is on par with some large pharma companies



hiPSC-derived cells produced in 2022, we believe that we are one of the largest hiPSCderived cell producers



## **ML-Based Analysis**

**Top 500 supercomputer across** any industry (TOP500 List, Nov 2022), we leverage vast neural networks and multiomics approaches to extract features and drive insights

#### **Recursion OS**

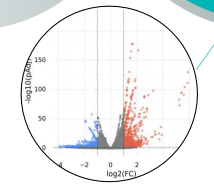
**Enables quality, relatability** and scale of data

## **High-Dimensional Validation**

Up to

15K

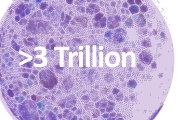
near whole exomes per week, we believe we are one of the largest transcriptomics data producers



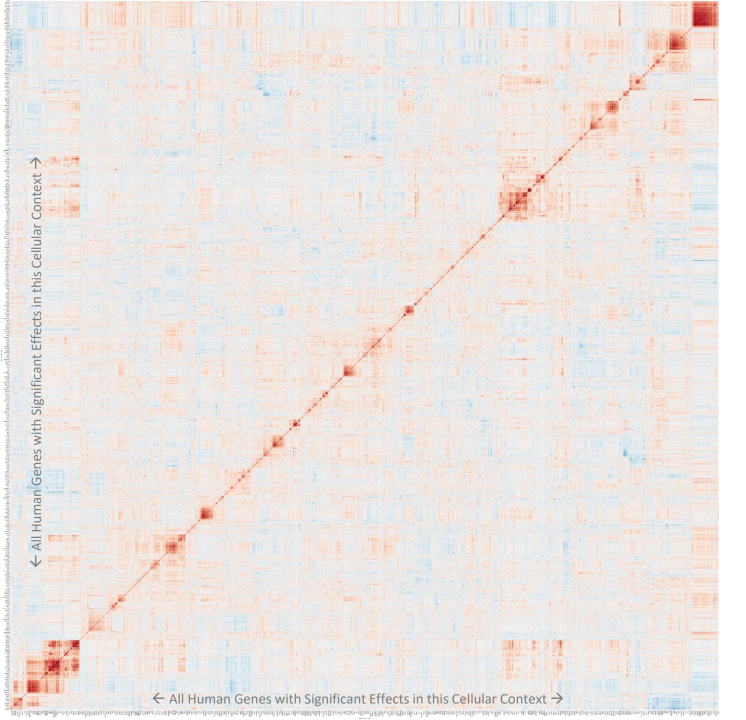
## **ML-Based Relationships**

Novel Insights at Scale

relatable hypotheses across multiple biological and chemical contexts







## **Genome-scale mapping**

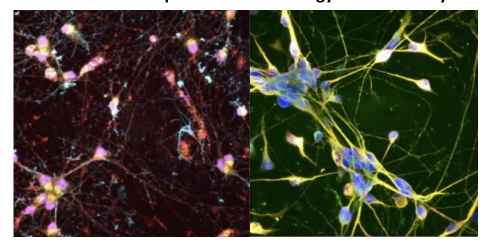
This is a **whole-genome arrayed CRISPR knock-out Map** generated in primary human endothelial cells

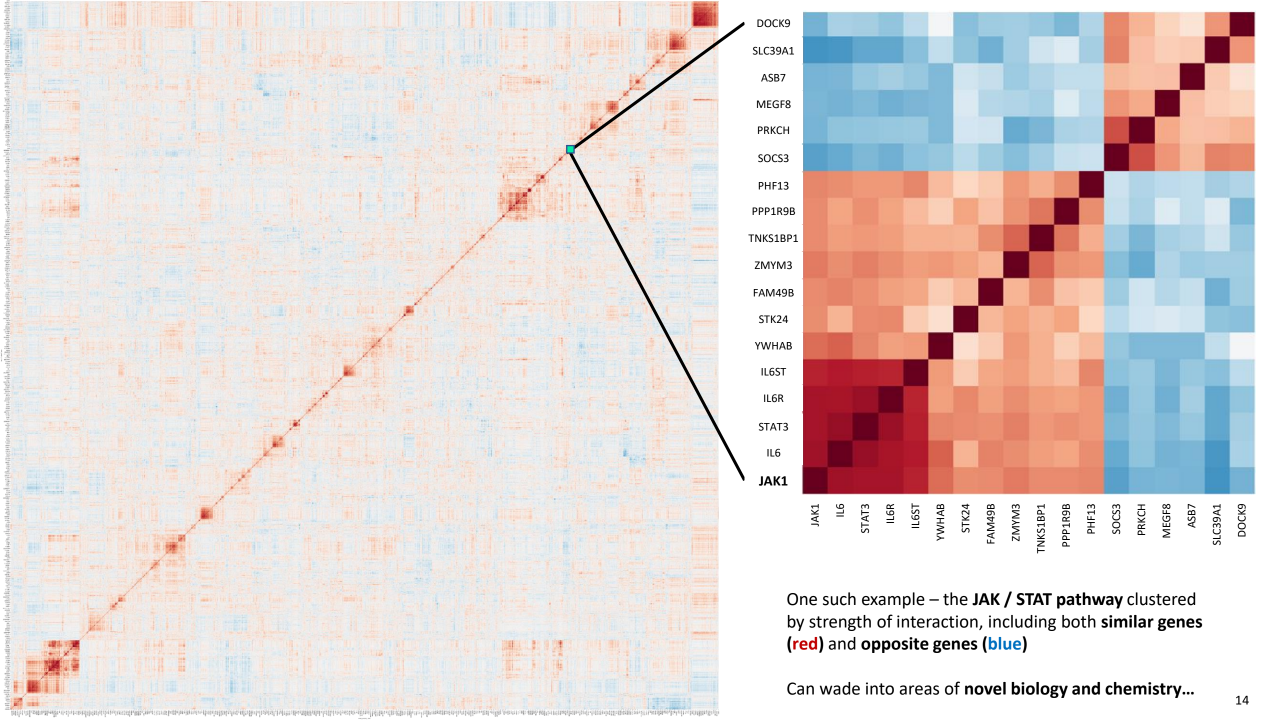
Every gene is represented in a pairwise way (each is present in columns and rows)

**Dark Red** indicates phenotypic similarity according to our neural networks while **Dark Blue** indicates phenotypic antisimilarity (which in our experience often suggests negative regulation)

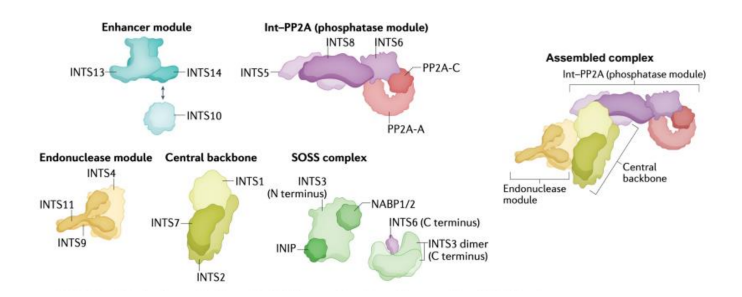
We can add the phenotypes of hundreds of thousands of small molecules at multiple doses and query and interact with these maps using a web application

#### Thousands of examples of known biology and chemistry

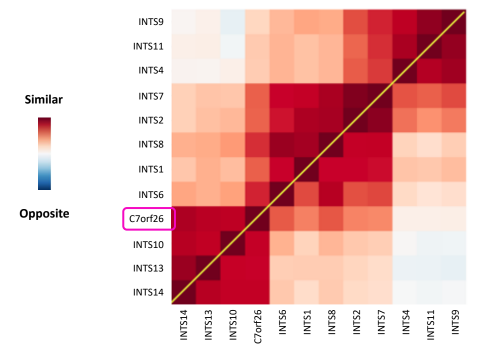








#### Phenomics TVN (below diagram) vs. Centerscale (above diagram)

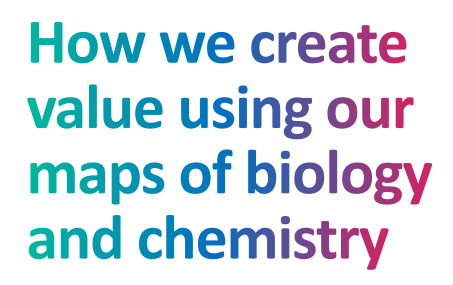


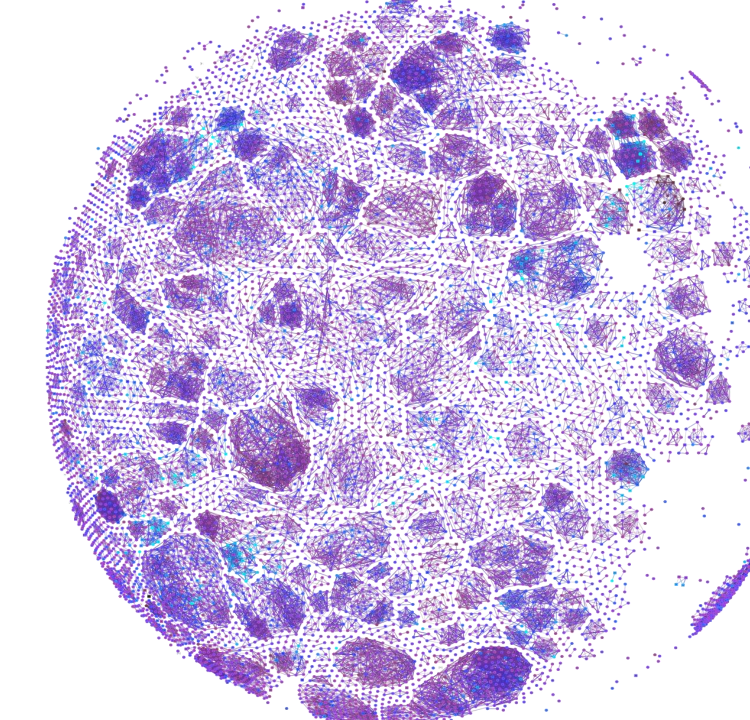
# Maps reveal known and novel biology

- In 2022, new independent research identified a previously unknown gene, C7orf26, as part of the Integrator complex
- Maps jointly developed by Recursion and Genentech replicated this same result
- Demonstrates accuracy and consistency across different map building approaches



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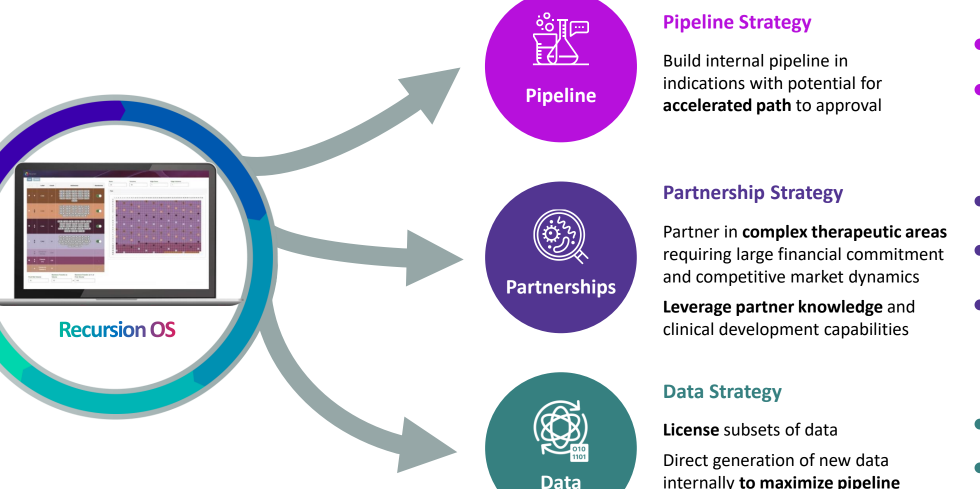






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## Harnessing value with a capital efficient business strategy



- **Precision Oncology**
- **Rare Disease**

- **Fibrosis**
- Neuroscience\*
- Other large, intractable areas of biology

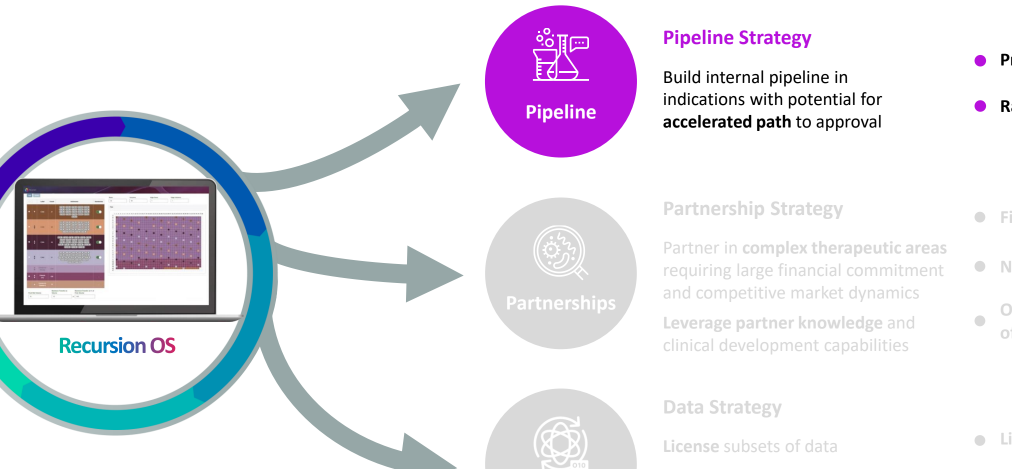
internally to maximize pipeline and partnership value-drivers

- Licensing
- **Augment Recursion OS**

\*Includes a single oncology indication from our Roche and Genentech collaboration.

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## Harnessing value with a capital efficient business strategy



- Precision Oncology
- Rare Disease

- Fibrosis
- Neuroscience\*
- Other large, intractable areas of biology

Direct generation of new data internally to maximize pipeline and partnership value-drivers

- Licensing
- Augment Recursion OS

ncludes a single oncology indication from our Roche and Genentech collaboration.



## Our pipeline reflects the scale and breadth of our approach

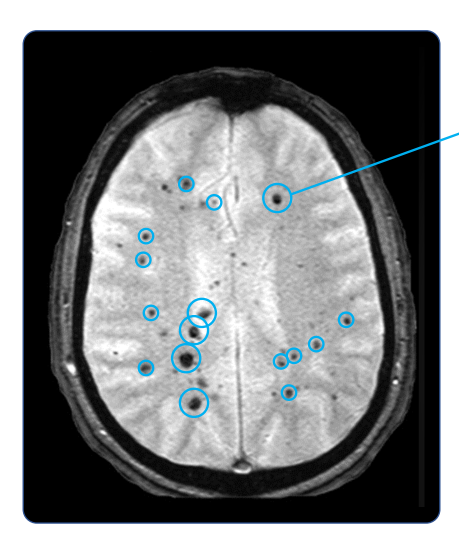


More than a dozen early discovery and research programs in oncology, neuroscience, inflammation & immunology, and rare disease

# **REC-994** for the Treatment of Symptomatic Cerebral Cavernous Malformations (CCM)

Target / MOA	Superoxide Scavenger
Molecule Type	Small Molecule
Lead Indication(s)	Cerebral Cavernous Malformations
Status	Phase 2
Designation(s)	US & EU Orphan Drug
Source of Insight	Recursion OS

## **Disease Overview: Cerebral Cavernous Malformations (CCM)**



#### Description

- Large unmet need for a novel nonsurgical treatment
- Vascular malformations (cavernomas) in the brain and spinal cord
- High-risk for hemorrhage creates "ticking time bomb"
- Progressive increase in CCM size and number over time in those with familial disease
- Debilitating symptoms, including intractable seizure, intracerebral hemorrhage, focal neurological deficits

"Historically, cavernomas have been managed primarily with observation, surgical resection, and occasionally radiotherapy. However, for a number of reasons, many patients with cavernomas must endure a life with neurologic symptoms"

- Ryan Kellogg, MD, Investigator at the University of Virginia

## **Disease Overview: Cerebral Cavernous Malformations (CCM)**



Julia - living with CCM

#### **Patient Population – Large and Diagnosable**

- >1 million patients worldwide live with these lesions today
- Caused by loss of function mutation in one of three genes: CCM1 (60%), CCM2 (20%), and CCM3 (20%)
- Inherited autosomal dominant mutation in 30-40%; or sporadic
- US symptomatic population is more than 5 times larger than other rare diseases like
   Cystic Fibrosis (>31k patients) and Spinal Muscular Atrophy (>33k patients)

#### **No Approved Medical Therapy**

- No approved drugs for CCM and no other potential therapeutic in industry-sponsored clinical development
- Most patients receive **no treatment** or only **symptomatic therapy**
- Surgical resection or stereotactic radiosurgery not always feasible because of location of lesion and is not curative

~360,000

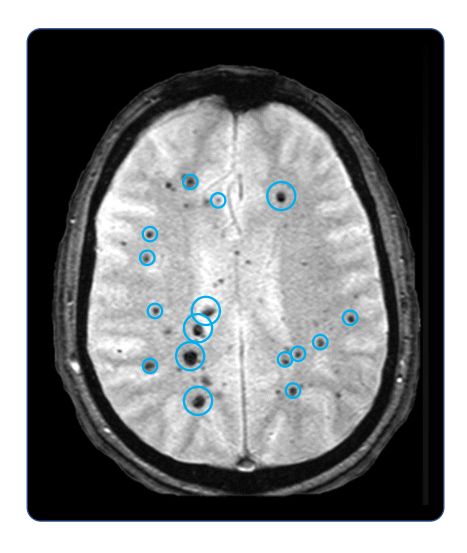
Symptomatic US + EU5 patients

## **Disease Overview: CCM is an Under-Appreciated Orphan Disease**

Non-oncology Orphan Indication	Product	U.S. + EU5 Prevalence	
Cerebral cavernous malformation (CCM)	REC-994 (Recursion)	>1,800,000 (Symptomatic: ~360,000)	
Idiopathic pulmonary fibrosis (IPF)	Esbriet (pirfenidone)	>160,000	
Cystic fibrosis (CF)	VX-669/ VX-445 + Tezacaftor + Ivacaftor - Vertex	>55,000	
Spinal muscular atrophy (SMA)	SPINRAZA (nusinersen)	>65,000	



## Therapeutic Approach to Cerebral Cavernous Malformations (CCM)

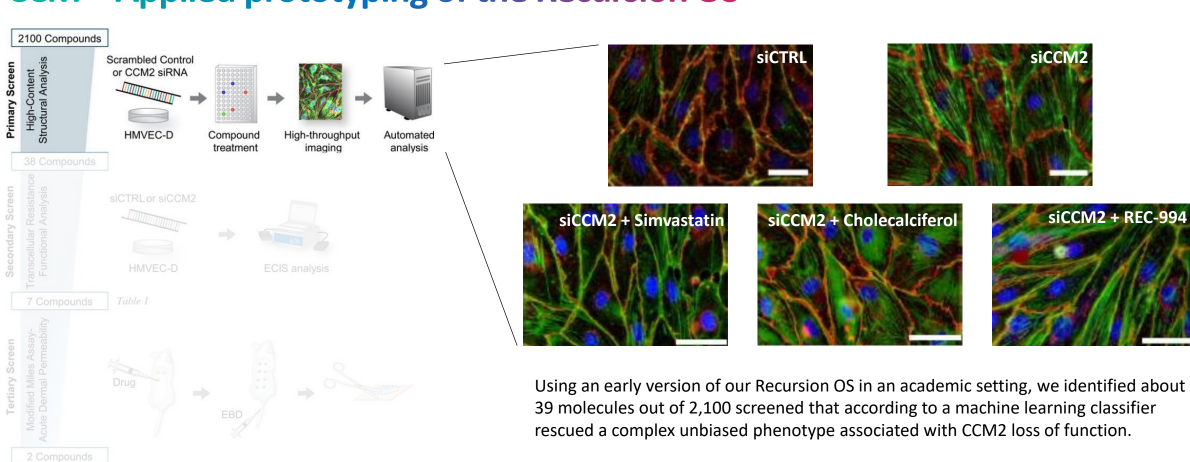


#### Novel therapeutic approach

- Symptoms associated with both increased size of lesions, but also inflammation or activation of lesions within the immunopriviledged environment of the brain
- Lesions arise from the capillary bed and are not high-pressure (e.g., the lesion growth is unlikely to be primarily driven by the *law of Laplace*)
- The Recursion Vascular Stability Hypothesis:
  - Eliminating the lesions may <u>not</u> be required for significant patient benefit
  - Slowing or halting the growth of the lesions while mitigating lesion leakiness and endothelial cell activation to halt the feed-forward inflammatory reaction may mitigate some symptoms and be beneficial to patients



## **CCM – Applied prototyping of the Recursion OS**

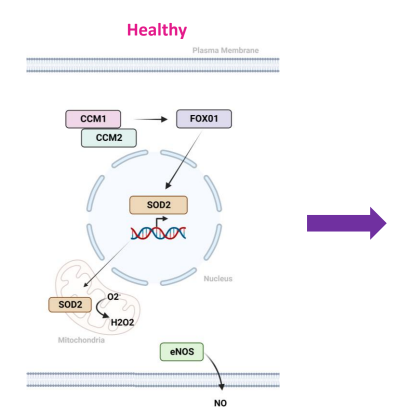


Through a set of follow-on confirmatory assays of increasing complexity, REC-994 stood out as one of two compounds we tested in a 5-month chronic CCM animal model where both compounds demonstrated significant benefit.



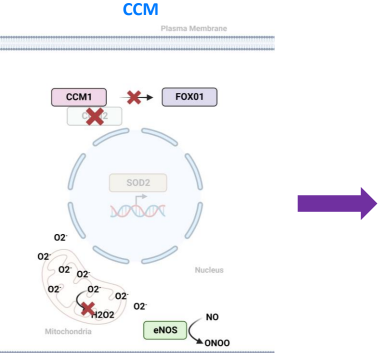
### Recursion

## **REC-994 – Mechanism of Action**



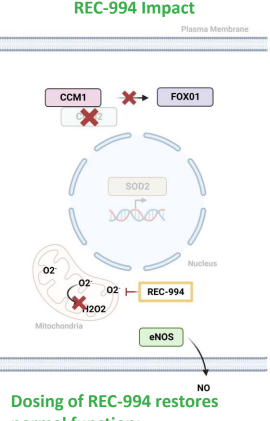
By regulating SOD2, CCM1 (KRIT1) & CCM2 suppress:

- Endothelial cell activation
- Smooth muscle proliferation
- Leukocyte adhesion
- Platelet aggregation



## **CCM1** or **CCM2** loss of function leads to activated endothelium:

- Decreased cell-cell junctional integrity and increased monolayer permeability
- Impaired vasodilation
- Cavernous angioma formation



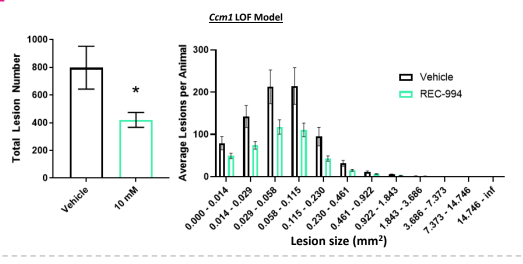
## normal function:

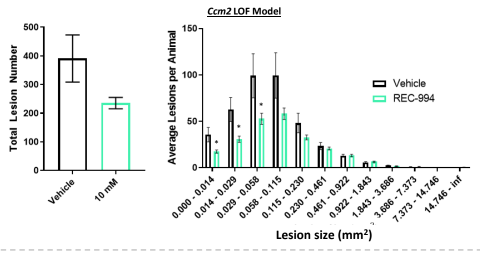
- Normalized ROS balance
- Restores quiescent endothelial cell state
- Stabilizes endothelial barrier function

## **Further Confidence: Preclinical Studies Confirm Insight**

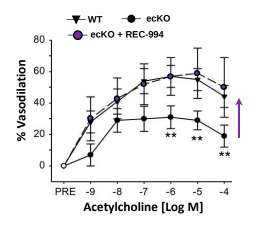
Preclinical Studies: REC-994 reduces lesion burden and ameliorates vascular defects in genetic mouse models of CCM

1 Reduces lesion number and size in *Ccm1* and *Ccm2* LOF mouse models



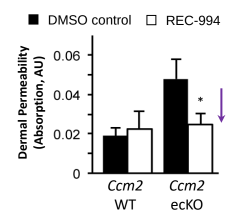


Completely rescues acetylcholine-induced vasodilation defect



REC-994 stabilizes the integrity of vasculature against challenges to permeability

#### Rescues dermal permeability defect in CCM2 mice



Vascular permeability is a clinically relevant feature of CCM lesions

## **Further Confidence : Clinical Studies Confirming Safety**

REC-994 Phase 1 Studies - well-tolerated with no dose-dependent adverse events in SAD and MAD

MAD Study	Placebo	50 mg	<b>200 mg</b>	400 mg	800 mg
Total Number of TEAEs	5	0	10	4	15
Total Subjects with ≥ one TEAE	4	0	3	3	4
Severity					
Mild	3	0	3	3	3
Moderate	1	0	0	0	1
Severe	0	0	0	0	0
Relationship to Study Drug					
None	3	0	0	2	1
Unlikely	1	0	1	1	2
Possibly	0	0	0	0	0
Likely	0	0	2	0	1
Definitely	0	0	0	0	0
Total Number of SAEs	0	0	0	0	0
Total Subject with ≥ one TEAE	0	0	0	0	0
Discontinued Study Drug Due to AE	0	0	0	0	0



## **SYCAMORE Clinical Trial: REC-994 Phase 2 Underway**

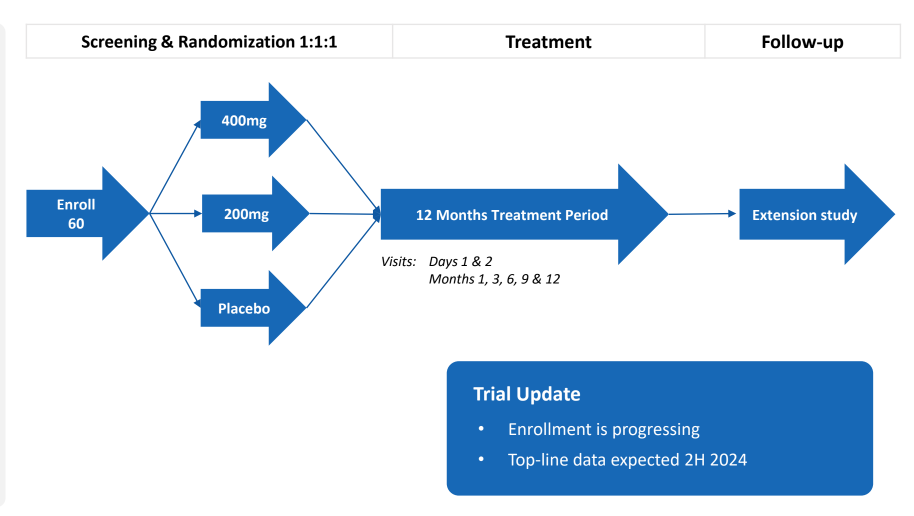
#### Phase 2 trial initiated in Q1 2022

#### **Enrollment Criteria**

- MRI-confirmed CCM lesion(s)
- Familial or sporadic
- Symptoms directly related to CCM

#### **Outcome Measures**

- Primary: Safety and tolerability
  - Adverse events & symptoms
- Secondary: Efficacy
  - Clinician-measured outcomes (CGI and PGI)
  - Imaging of CCM lesions number, size & rate of change
  - Impact of acute stroke (mRS, NIHSS)
  - Patient reported outcomes (SMSS, PROMIS-29, CCM HI, symptom questionnaires)
- Exploratory: Biomarkers



# **REC-2282** for the Treatment of Progressive Neurofibromatosis Type 2 (NF2) Mutated Meningiomas

Target / MOA	HDAC Inhibitor
Molecule Type	Small Molecule
Lead Indication(s)	NF2 Mutated Meningiomas
Status	Phase 2/3
Designation(s)	Fast Track; US and EU Orphan Drug
Source of Insight	Recursion OS

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#### Clinical: NF2

## **Disease Overview: Neurofibromatosis Type 2 (NF2)**



Ricki – living with NF2

### **Patient Population – Large and Diagnosable**

- Rare autosomal dominant tumor syndrome resulting from biallelic inactivation of the NF2 gene which leads to deficiencies in the tumor suppressor protein merlin
- NF2 can be inherited or spontaneous (>50% of patients represent new mutations);
   up to 1/3 are mosaic
- CNS manifestations: meningiomas and vestibular schwannomas; mean age at presentation: ~20 years

## **No Approved Medical Therapy**

- No approved drugs for NF2
- Surgery is standard of care (when feasible)
- Location may make complete resection untenable, leading to hearing loss, facial paralysis, poor balance and visual difficulty

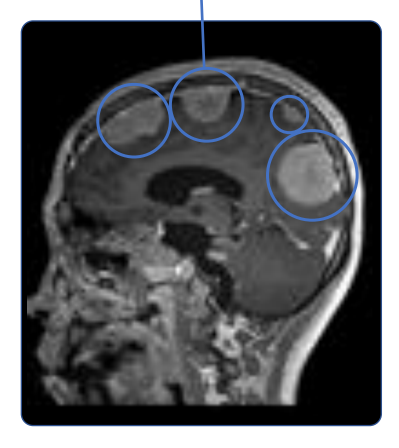
Source: https://rarediseases.org/rare-diseases/neurofibromatosis-2

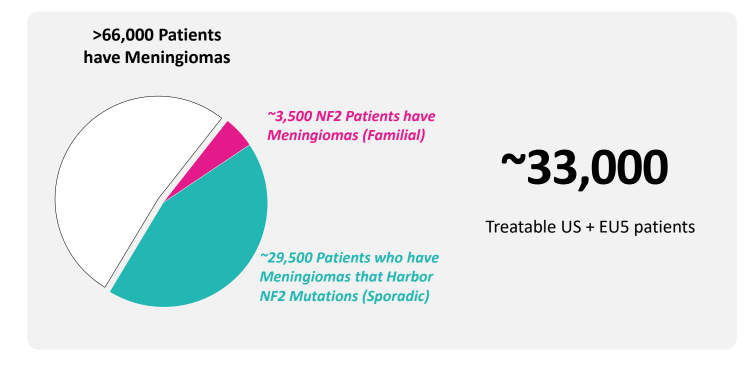


## Disease Overview: Neurofibromatosis Type 2 (NF2) Meningiomas

- Most tumors are benign and slow growing but location in CNS leads to serious morbidity or mortality
- Prognosis is adversely affected by early age at onset, a higher number of meningiomas and having a truncating mutation

## **Intracranial Meningioma**

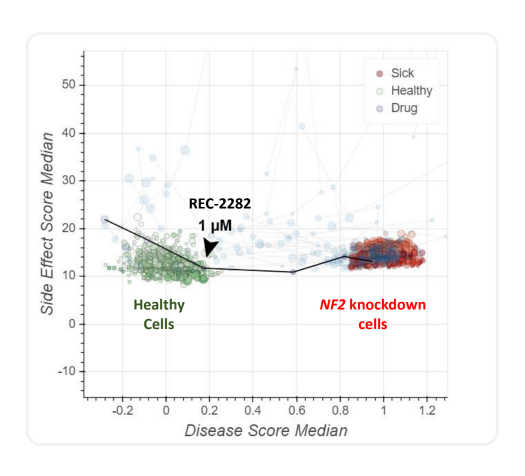




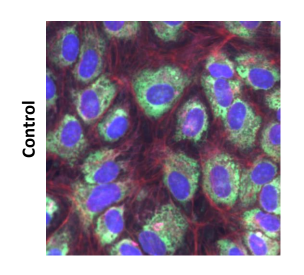
- Threatens mortality; if amenable, surgical excision is primary intervention
- Many patients have multiple meningiomas that exhibit heterogenous behavior and asynchronous growth
- Stasis or shrinkage of tumor could improve prognosis

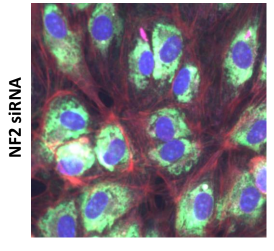


## Insight from OS: REC-2282 Rescued Loss of NF2



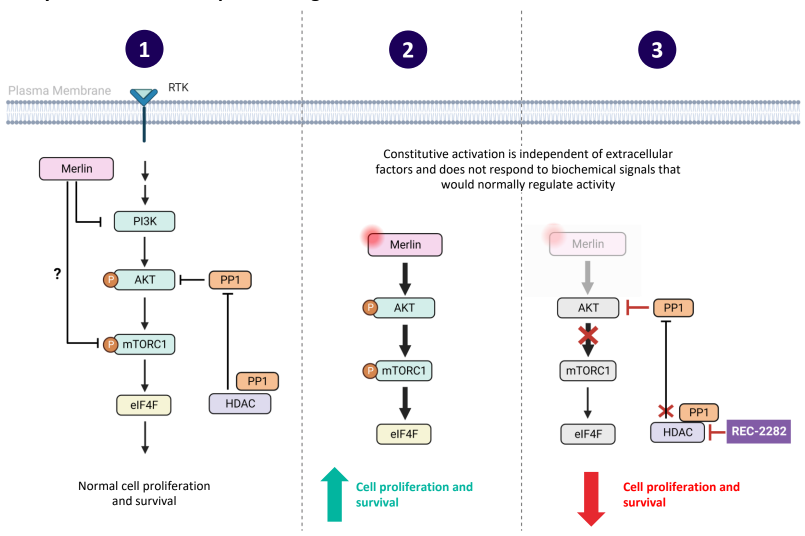
## REC-2282 identified as rescuing HUVEC cells treated with NF2





## **REC-2282 – Mechanism of Action**

#### Orally Bioavailable, CNS-penetrating, Small Molecule HDAC Inhibitor



NF2 encodes for the protein Merlin and negatively regulates mTOR signaling

Loss of Merlin leads to increased signaling in the PI3K/AKT/mTOR pathway

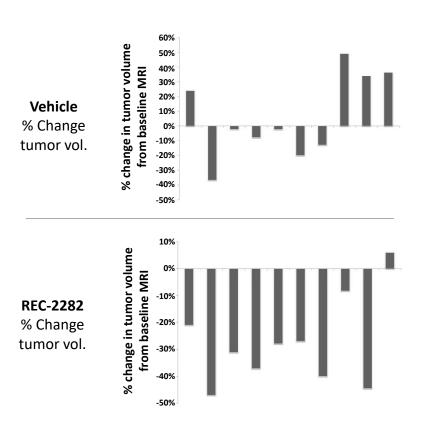
Oncogenic mTOR signaling arrested with HDAC inhibitors



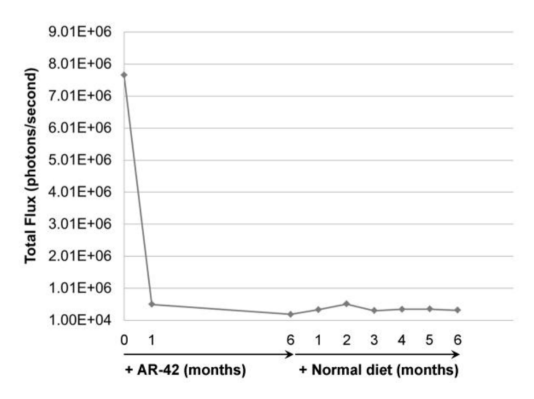
## **Further Confidence: Preclinical Studies Confirming Insight**

REC-2282 preclinical studies demonstrated clear in-vivo efficacy in multiple NF2 tumor types

1 Shrinks vestibular schwannoma xenografts in nude mice



2 Prevents growth & regrowth of NF2-deficient meningioma model in mice





## **Further Confidence: Prior Studies Suggest Potential Therapeutic Benefit**

- Evaluable Patients: CNS Solid Tumors: NF2 N=5; Non-CNS Solid Tumors: N=10
- PFS: CNS solid tumors = 9.1 months; Non-CNS solid tumors =
   1.7 months
- Best overall response = SD in 8/15 patients (53%; 95% CI 26.6–78.7)
- Longest duration of follow-up without progression: > 27
  months (N=1)
- Most common AEs: cytopenia, fatigue, nausea



## Well understood clinical safety ...



Multiple investigator-initiated studies in oncology indications



Lengthy human clinical exposure in NF2 – multiple patients on drug for several years



Well-characterized side effect profile

## ... with a drug-like profile



Established and scalable API manufacturing process



Multiple cGMP batches of 10mg and 50mg tablets have been manufactured

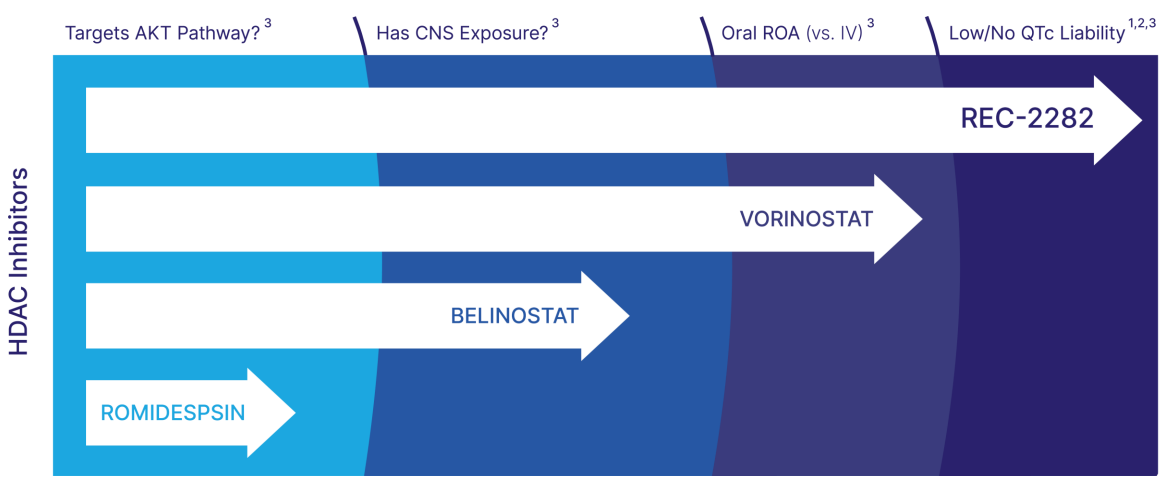


**Excellent long-term stability** 

Clinical: NF2

## **REC-2282 Appears Well Suited for NF2 vs Other HDAC Inhibitors**

#### **REC-2282 Would be First-In-Class HDAC Inhibitor for Treatment of NF2 Meningiomas**



<sup>&</sup>lt;sup>1</sup>Sborov DW, et al. A phase 1 trial of the HDAC inhibitor AR-42 in patients with multiple myeloma and T- and B-cell lymphomas. Leuk Lymphoma. 2017 Oct;58(10):2310-2318.

<sup>&</sup>lt;sup>2</sup>Collier KA, et al. A phase 1 trial of the histone deacetylase inhibitor AR-42 in patients with neurofibromatosis type 2-associated tumors and advanced solid malignancies. Cancer Chemother Pharmacol. 2021 May;87(5):599-611.

<sup>&</sup>lt;sup>3</sup> Prescribing Information of Vorinostat/Belinostat/Romidepsin respectively



## POPLAR Clinical Trial: REC-2282 for NF2 Phase 2/3 Underway

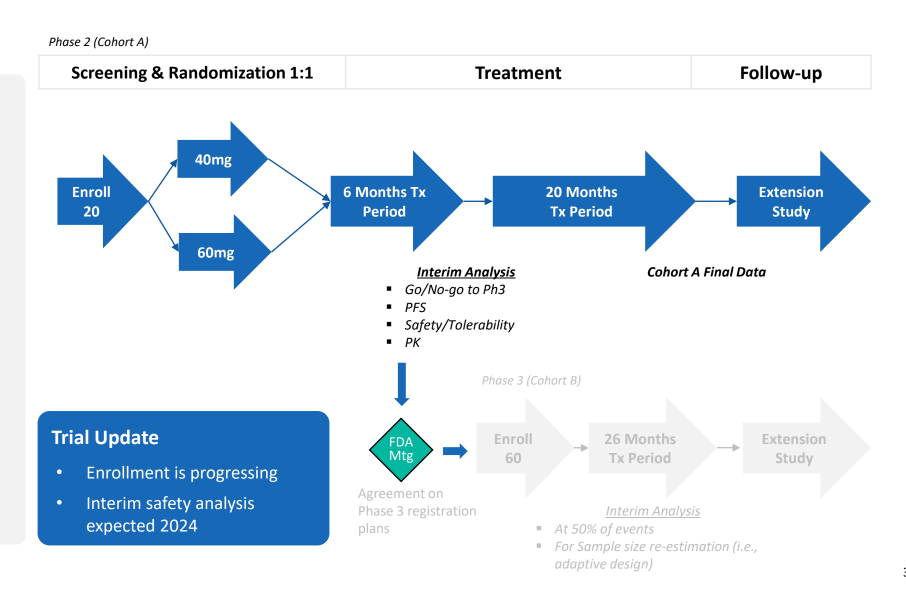
#### Phase 2/3 trial initiated in Q2 2022

#### **Enrollment Criteria**

- MRI-confirmed progressive meningioma
- Either of the below
  - Sporadic meningioma with confirmed NF2 mutation
  - Confirmed diagnosis of NF2 disease

#### **Outcome Measures**

- Primary: Safety and tolerability
  - Progression-free survival
  - Time to progression
  - Duration of response
  - Overall response rate



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# **REC-4881** for the Treatment of Familial Adenomatous Polyposis (FAP)

Target / MOA	MEK Inhibitor	
Molecule Type	Small Molecule	
Lead Indication(s)	Familial Adenomatous Polyposis	
Status	Phase 2	
Designation(s)	Fast Track; US and EU Orphan Drug	
Designation(s)  Source of Insight	Fast Track; US and EU Orphan Drug  Recursion OS	



## **Disease Overview: Familial Adenomatous Polyposis**



Polyps Found in Colon and Upper GI Tract

#### **Patient Population – Easily Identifiable**

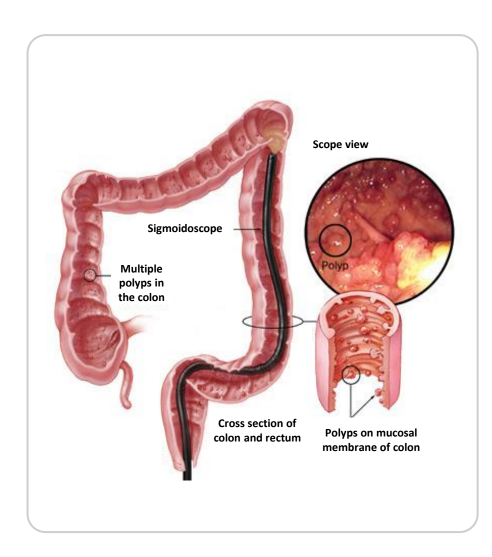
- Autosomal dominant tumor predisposition syndrome caused by a mutation in the APC gene
- Classic FAP (germline mutation) :
  - Hundreds to thousands of polyps in colon and upper GI tract
  - Extraintestinal manifestations (e.g., desmoid tumors)
  - 100% likelihood of developing colorectal cancer (CRC) before age 40, if untreated

~50,000

Diagnosed US + EU5 patients



## Disease Overview: Familial Adenomatous Polyposis – Standard of Care



#### **No Approved Medical Therapy**

- Standard of care: colectomy during adolescence (with or without removal of rectum)
- Post-colectomy, patients still at significant risk of polyps progressing to GI cancer
- Significant decrease in quality-of-life post-colectomy: continued endoscopies and surgical intervention

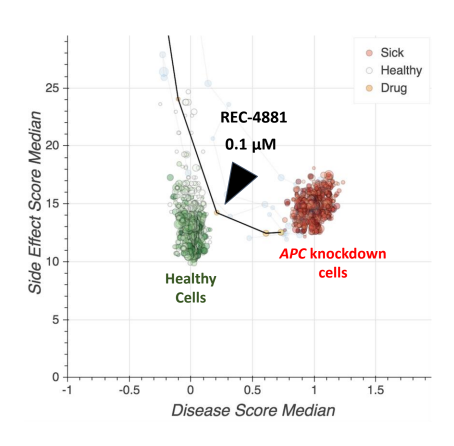
"Despite progress with surgical management, the need for effective therapies for FAP remains high due to continued risk of tumors post-surgery"

- Niloy Jewel Samadder, MD, Mayo Clinic



## Insight from OS: Rescued Loss of APC, Inhibited Tumor Growth

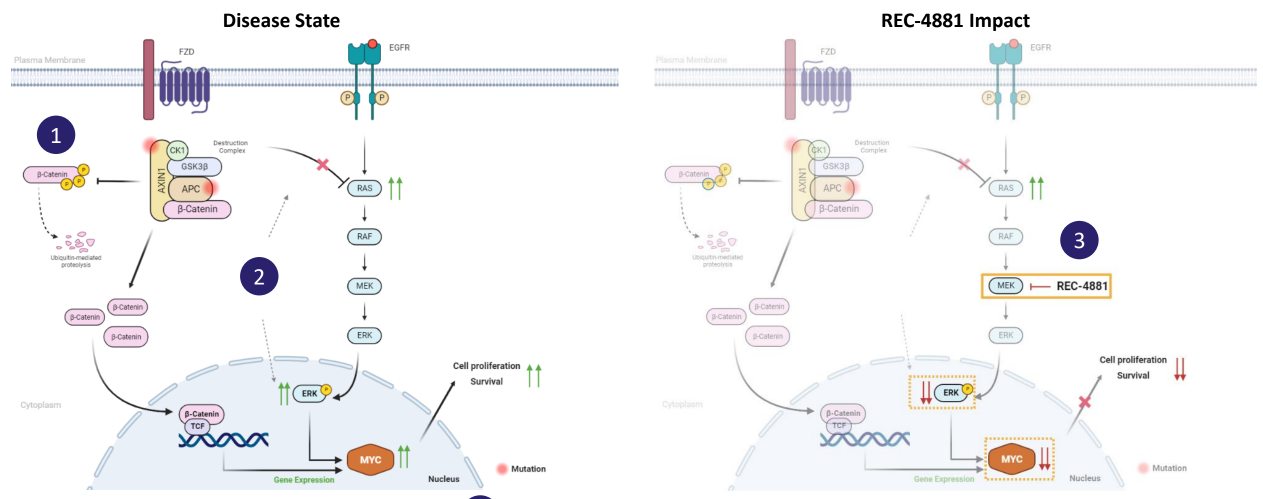
## REC-4881 rescued phenotypic defects of cells with APC knockdown



- Compared to thousands of other molecules tested, REC-4881 rescued phenotypic defects substantially better (including better rescue than other MEK inhibitors) for APC specific knockdown
- Findings validated in tumor cell lines and spheroids grown from human epithelial tumor cells with APC mutation
  - 1,000x more selectivity in tumor cell lines with APC mutation
  - Inhibited growth and organization of spheroids

## MoA: REC-4881 Blocks Wnt Mutation Induced MAPK Signaling

#### **Orally Bioavailable, Small Molecule MEK Inhibitor**



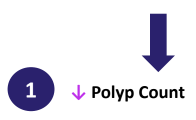
REC-4881 inhibits MEK 1/2 and recovers the destabilization of RAS by the β-Catenin destruction complex, restoring the cell back to a Wnt-off like state



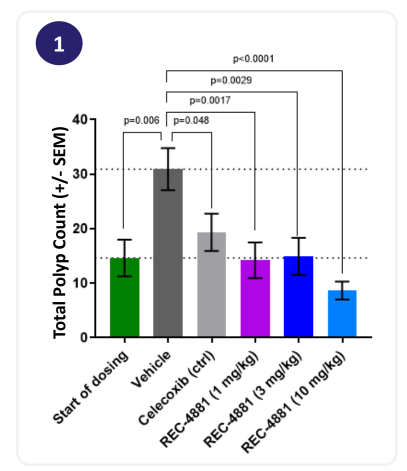
# Further Confidence: Preclinical Studies Confirming Reduction in Polyp Count and High-Grade Dysplasia

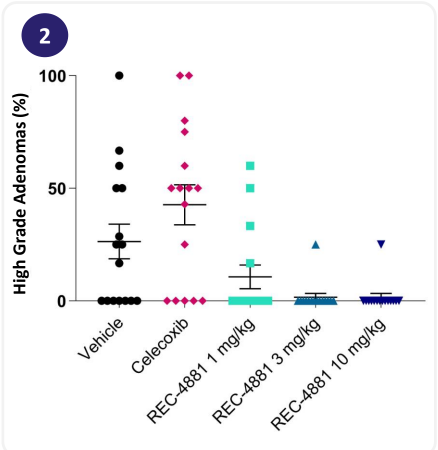
- In-vivo efficacy in APCmin mouse model
- Apc<sup>min</sup> = FAP disease model
- Mice treated once daily for 8 weeks

After 8 weeks of treatment:









## **Further Confidence: Clinical Data Generated by Recursion**

REC-4881-101: Single-center, double-blind, placebocontrolled, dose-escalation study in healthy volunteers

- Group 1 (n=13): Food effect crossover (REC-4881 4 mg/PBO [fed/fasted]), followed by single dose REC-4881 8 mg/PBO [fed]
- Group 2 (n=12): Matched single ascending dose (REC-4881 4 mg/PBO; REC-4881 8 mg/PBO; REC-4881 12 mg/PBO)

#### Accomplished



Recursion formulation yields exposures comparable to Takeda's formulation (molecule in-licensed from Takeda)



No food effect



Dose proportional increases in exposure



Similar to C20001 study, observed pERK inhibition (i.e., target engagement) at 8 mg and 12 mg doses



Acceptable safety profile





## **TUPELO Clinical Trial: REC-4881 for FAP Phase 2 Underway**

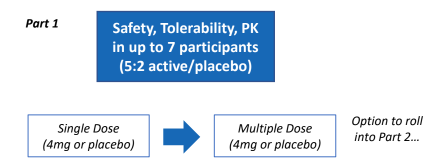
#### Phase 2 trial initiated in Q3 2022

#### **Enrollment Criteria**

- Confirmed APC mutation
- Post-colectomy/proctocolectomy
- No GI cancer present
- Polyps in either duodenum (including ampulla of vater) or rectum/pouch

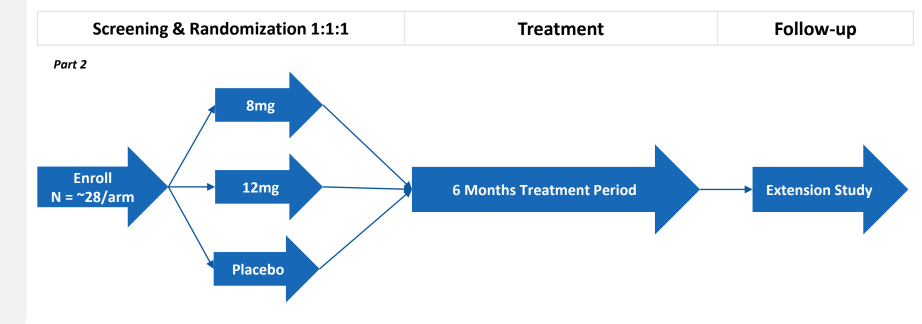
#### **Outcome Measures**

- Primary:
  - Part 1: PK
  - Part 2: % change from baseline in polyp burden
- Secondary:
  - Part 1: Safety & tolerability
  - Part 2: PK; PD; change from baseline in polyp number, histological grade, disease scoring
- Exploratory:
  - Part 1: PD
  - Part 2: Time to first occurrence of FAPrelated event; change from baseline in extent of desmoid disease



#### **Trial Update**

 Recent protocol amendments aimed at accelerating quality and pace of the trial



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# **REC-4881** for the Treatment of Solid Tumors with AXIN1 or APC Mutant Cancers

Target / MOA	MEK Inhibitor	
Molecule Type	Small Molecule	
Lead Indication(s)	Solid Tumors with AXIN1 or APC Mutant Cancers	
Status	Phase 1b/2	
Source of Insight	Recursion OS	

### Disease Overview: AXIN1 or APC Mutant Cancers



Gross morphology of HCC tumor

- Sustained Wnt signaling is a frequent driver event found across a wide variety of solid tumors
- Dysregulation of  $\beta$ -catenin destruction complex due to inactivating mutations in *AXIN1* or *APC* leads to sustained Wnt signaling promoting cancer progression and survival<sup>1</sup>
- AXIN1 or APC mutant solid tumors are considered clinically aggressive and resistant to standard treatments

"Nothing in HCC has immediate therapeutic relevance and the most common mutations are TERT, TP53, and Wnt (CTNNB1/AXIN1/APC) and combined these alterations define almost 80% of patients and are not targetable"

- KOL, Clinical Investigator, Texas

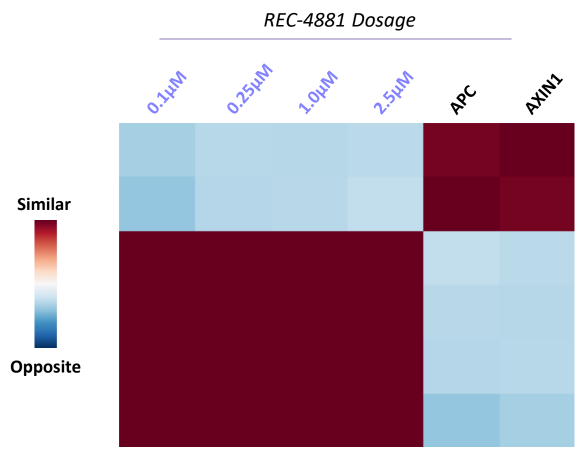
<sup>1</sup>Bugter, J.M., et al. Nat Rev Cancer, 2021, 21, pp.5-21

## Disease Overview: AXIN1 or APC Mutant Cancers

Tumor	AXIN1	APC	Treatable	Flexible Patient Selection Strategy and Study Design
Туре	Mutation Frequency <sup>1</sup>	Mutation Frequency <sup>1</sup>	Population <sup>2</sup> (US+EU5)	<ul> <li>AXIN1 and APC genes covered by commercially available NGS</li> </ul>
CRC	3%	70%	27,450	panels and liquid biopsy detection assays
LUAD	4%	11%	14,000	FDA guidance supports utility of ctDNA as patient selection for
Prostate	2%	11%	6,700	the detection of alterations for eligibility criteria and as a stratification factor for trials enrolling marker-positive and
Bladder	3%	8%	5,100	marker-negative populations <sup>3</sup>
НСС	12%	5%	3,100 —	Multiple tumor types will inform study design and patient
Endometrial	8%	12%	2,600	selection
Esophageal	2%	7%	2,600	
PDAC	1%	2%	1,500	Preclinical data with REC-4881 at clinically relevant
Ovarian	1%	3%	1,400 —	exposures in HCC and Ovarian PDX mouse models gives confidence to pursue other mutant
TNBC	1%	2%	300	cancer types
			~65,000	



## **Insight from OS: Novel Insight around Established MoA**



Heat map from Recursion OS

**Hypothesis:** Rescue of *AXIN1* may impact tumor progression and/or restore checkpoint sensitivity in cancers driven by *AXIN1* loss

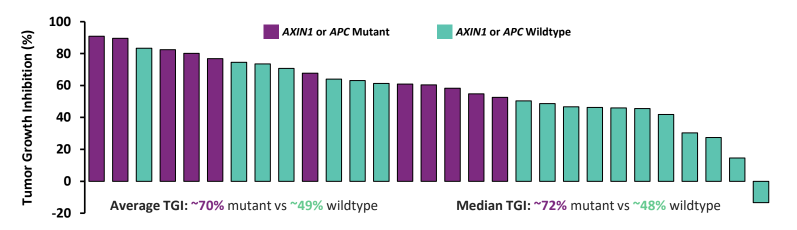
**Recursion Differentiation:** REC-4881 rescues tumor suppressor genes *APC* and *AXIN1* 

- APC and AXIN1 are negative regulators of Wnt signaling
- Both proteins form part of the B-catenin destruction complex. Strong clustering suggests map recapitulation of this biology

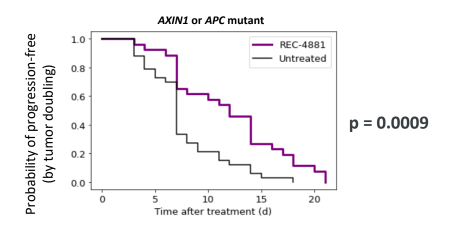


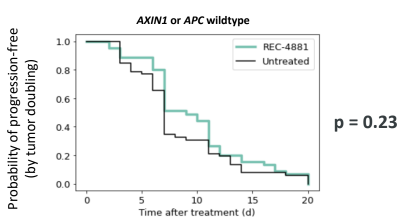
## **Further Confidence : Preclinical Studies Confirming Insight**

#### Efficacy found in In Vivo Mice Models ...



#### ... Led to Significant Progression Free Survival





#### **Next Steps**

- Finalize design of a Phase 1b/2 biomarker-enriched trial
- □ Initiate Phase 1b/2 trial in select tumor types in early 2024
- Identify suitable partners for genetic testing capabilities
- Evaluate REC-4881 in combination with targeted and/or immune modulating agents

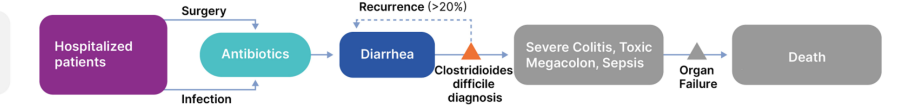
# **REC-3964** for the Treatment of C. Difficile Infection

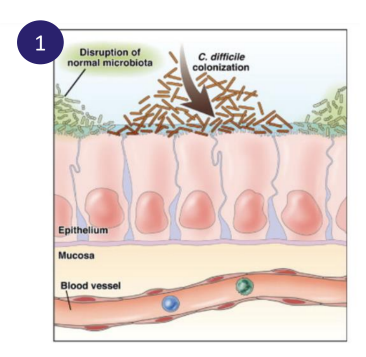
Target / MOA	Selective C. diff Toxin Inhibitor	
Molecule Type	Small Molecule	
Lead Indication(s)	ndication(s) C. Difficile Infection	
Status	Phase 1	
Source of Insight	Recursion OS	



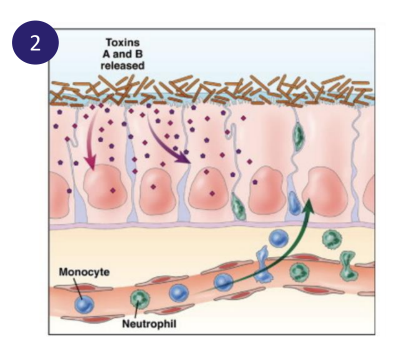
## **Disease Overview: C. Difficile Infection (CDI)**

C.diff is the leading cause of antibiotic-associated diarrhea

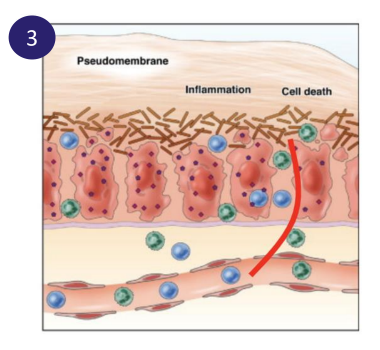




Disruption of microbiota and colonization of *C. diff* 



Release of C. diff toxins



Degradation of colon cell junction & toxin transit to bloodstream

## **Disease Overview : C. Difficile Infection (CDI)**



Colleen – lived with rCDI

#### Patient Population – Large, Diagnosable and Easy to Identify

- Symptoms caused by clostridioides difficile tissue-damaging toxins released in the colon
- Patients who experience >3 unformed stools are diagnosed via NAAT\* for toxin gene or positive stool test for toxins
- Patients who are at highest risk are those on antibiotics, and frequently visit hospitals or are living in a nursing home
- More than 80% of cases occur among patients age 65 or older

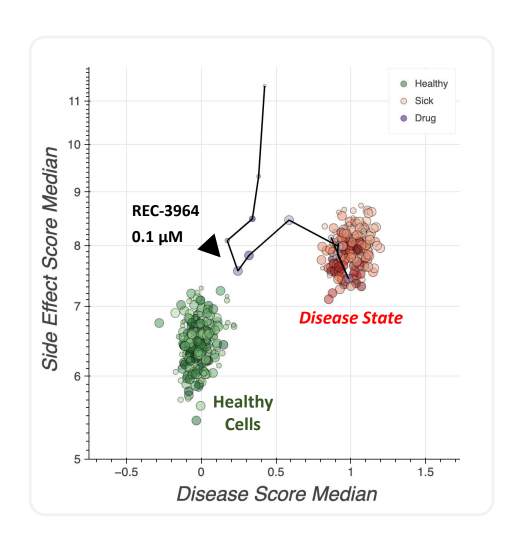
#### Large, Unmet Need with Significant Cost Burden

- RCDI\*\* occurs in 20-30% of patients treated with standard of care
  - 40% of those patients will continue to recur with 2+ episodes
- >29,000 patients die in the US each year from CDI
- Cost burden of up to \$4.8bn annually

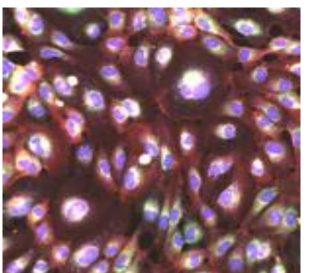
~730,000

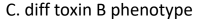
**Diagnosed US + EU5 patients** 

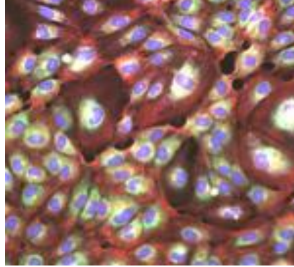
## **Insight from OS: REC-3964 Rescued Cells Treated with C. Difficile Toxins**



## REC-3964 identified as a NCE that demonstrated strong rescue in HUVEC cells treated with C. diff toxin



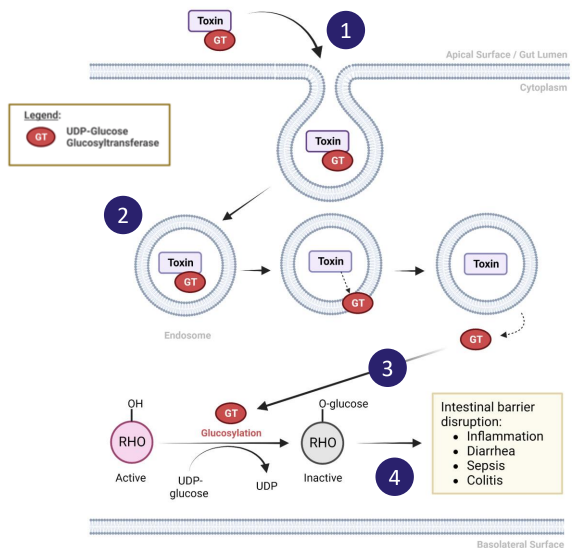




**Healthy Control** 

## **REC-3964: Selective Inhibitor of C. Difficile Toxins**

#### REC-3964 is Recursion's 1st Small Molecule NCE to Reach the Clinic

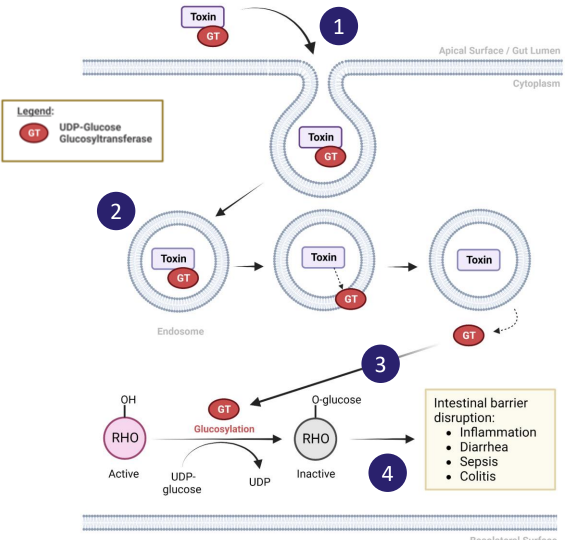


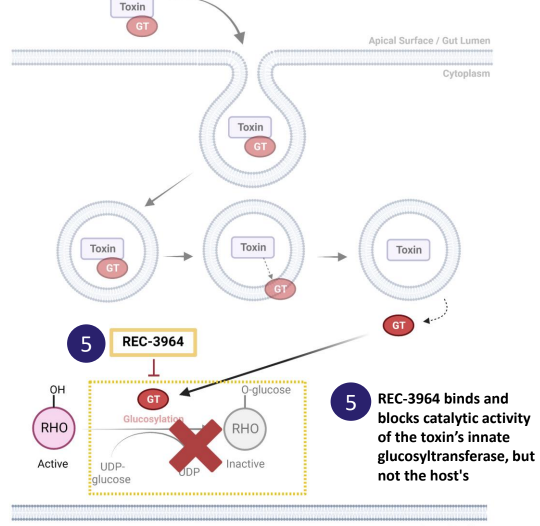
- C.diff toxins bind to cell surface receptors and trigger endocytic event
- Autocatalytic cleavage event releases C.diff toxin's glucoyltransferase enzymatic domain into the cytosol of the infected cell
- The glucosyltransferase locks Rho family GTPases in the inactive state
- Inactivation of Rho GTPases alters cytoskeletal dynamics, induces apoptosis, and impairs barrier function which drives the pathological effects of C.diff infection

Adapted from Awad et al. 2014

## **REC-3964: Selective Inhibitor of C. Difficile Toxins**

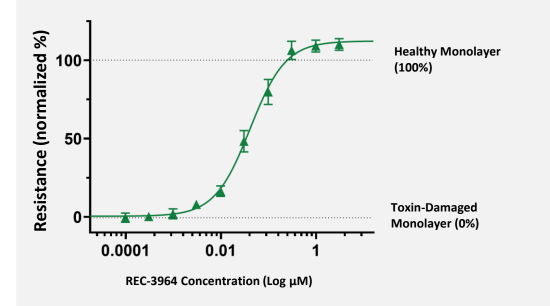
#### REC-3964 is Recursion's 1st Small Molecule NCE to Reach the Clinic





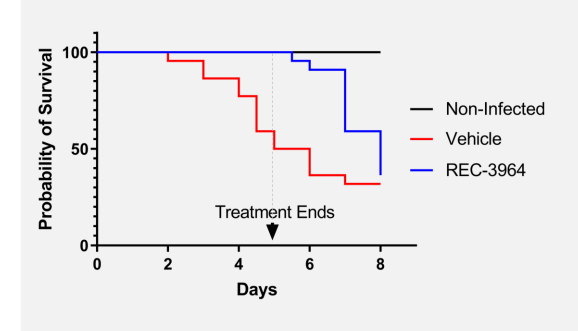
## **Further Confidence: Preclinical Studies Confirmed Recursion OS Insight**

## REC-3964 rescues barrier integrity with increasing concentrations



✓ REC-3964 restores gut epithelial barrier integrity, which when disrupted causes inflammation and diarrhea

## REC-3964 improved probability of survival in a hamster model of C. difficile infection



 Improved probability of survival beyond treatment completion



## Clinical Trial: REC-3964 for C. difficile Phase 1 Study Underway

#### Phase 1 FIH SAD/MAD Trial initiated in Q3 2022

#### **Trial Design**

Randomized, Double-blind Trial

#### **Population**

- Healthy Subjects
- SAD (n = 56)
- MAD (n = 50)

#### **Primary Objectives**

- Assess the safety & tolerability of SAD and MAD of REC-3964
- ✓ Evaluate the PK profile of REC-3964 after single and multiple doses

#### **Trial Update**

- Enrollment is progressing
- In several SAD cohorts and at least one MD cohort, REC-3964 has been extremely safe and well tolerated
- Complete safety and PK data readout expected
   2H 2023

## Preclinical Programs

RBM39: HR-Proficient Ovarian Cancer

Target α: Immunotherapy



# RBM39: Novel CDK12-Adjacent Target for Potentially Treating HR-Proficient Ovarian Cancer

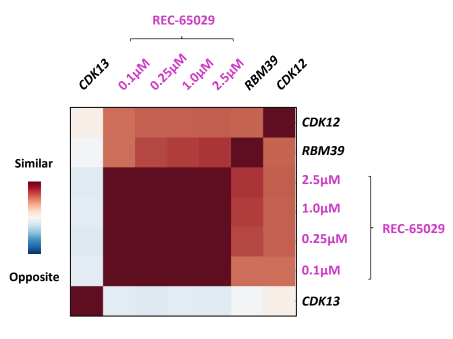
Identify potential first-in-class tumor-targeted precision therapeutic NCE with novel MOA capable of potentially treating HR-proficient ovarian cancer

Inhibition of target RBM39 (previously referred to as Target γ) may mimic the inhibition of CDK12 while mitigating toxicity related to CDK13 inhibition

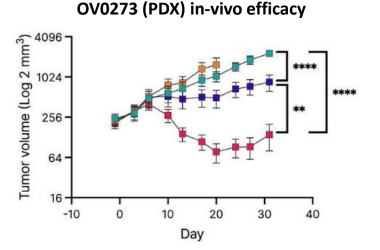
A Recursion-generated NCE showed single agent efficacy that is enhanced in combination with Niraparib in a BRCA-proficient PDX model

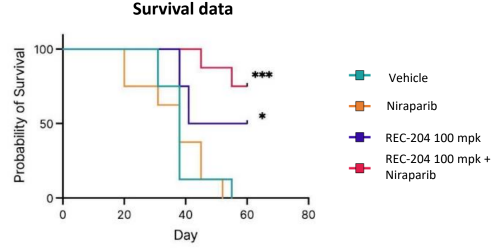
NEXT STEPS

Program anticipates reaching IND-enabling studies in 2023











## Target $\alpha$ : Potential First-in-Class NCE with **Novel MOA to Enhance Anti-PD-(L)1 Response**

**GOAL INSIGHT FROM OS**  Identify novel compounds capable of enhancing the therapeutic benefit of checkpoint therapy without concomitant inflammatory side effects

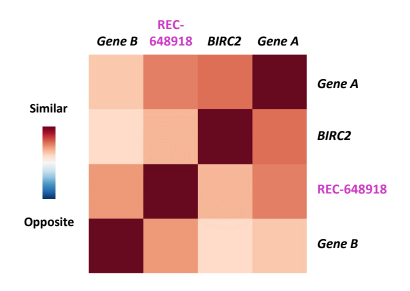
Novel compound identified with similarity to knockout of potential immunotherapy resistance gene targets (Gene A, Gene B)

**FURTHER CONFIDENCE** 

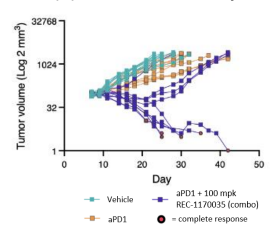
A Recursion-generated NCE showed reduction in tumor growth vs. anti-PD-1 alone in CT26 checkpoint resistance model - including 60% complete responses

**NEXT STEPS** 

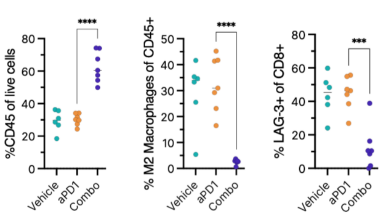
Program anticipates reaching IND-enabling studies in 2023



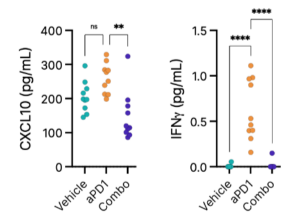
#### (A) CT26 in vivo efficacy



#### (B) Tumor Microenvironment Modulation



#### (C) Suppressed Peripheral Inflammation



## Harnessing value with a capital efficient business strategy





#### **Pipeline Strategy**

accelerated path to approval

- **Precision Oncology**
- **Rare Disease**



#### **Partnership Strategy**

Partner in complex therapeutic areas requiring large financial commitment and competitive market dynamics

Leverage partner knowledge and clinical development capabilities

- **Fibrosis**
- Neuroscience\*
- Other large, intractable areas of biology



#### **Data Strategy**

**License** subsets of data

Direct generation of new data internally to maximize pipeline and partnership value-drivers

- Licensing
- **Augment Recursion OS**

**Recursion OS** 



# Our existing partnerships represent some of the most significant scientific collaborations in biopharma



(Announced Sep 2020; Expanded Dec 2021)



- \$30M upfront and \$50M equity investment
- Up to or exceeding \$1.2B in milestones for up to or exceeding
   12 programs
- Mid single-digit royalties on net sales
- Recursion owns all algorithmic improvements



(Announced Dec 2021)

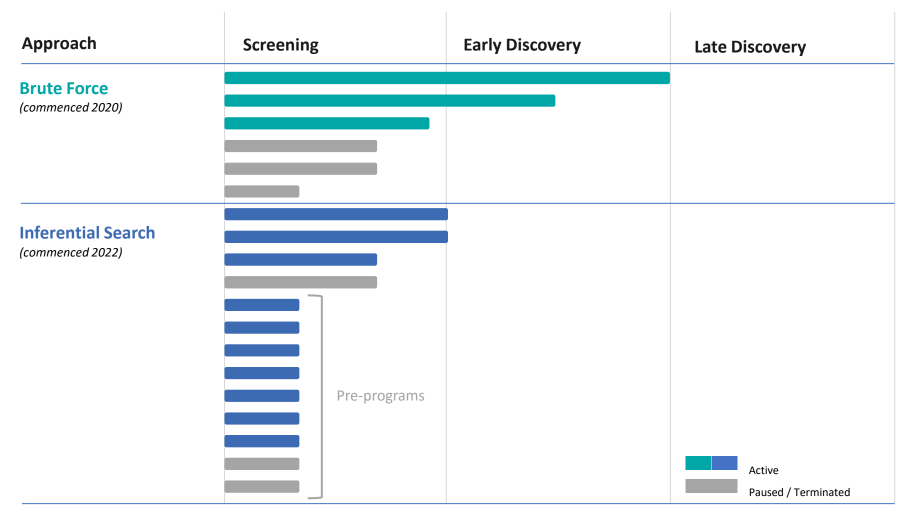


- \$150M upfront and up to or exceeding \$500M in research milestones and data usage options
- Up to or exceeding \$300M in possible milestones per program for up to 40 programs
- Mid to high single-digit tiered royalties on net sales
- Recursion owns or co-owns all algorithmic improvements



## Multiple programs advancing in parallel to near-term milestones

#### Transition to Inferential Search has accelerated new program initiation in 2022





# Our existing partnerships represent some of the most significant scientific collaborations in biopharma



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- \$30M upfront and \$50M equity investment
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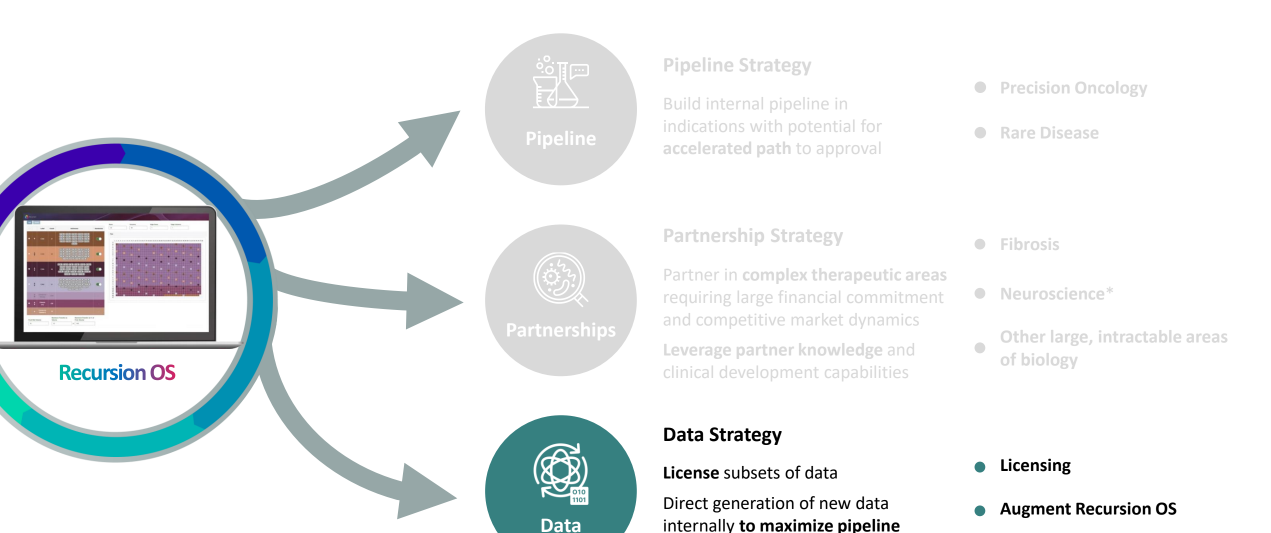
(Announced Dec 2021)



- \$150M upfront and up to or exceeding \$500M in research milestones and data usage options
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- Mid to high single-digit tiered royalties on net sales
- Recursion owns or co-owns all algorithmic improvements

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## Harnessing value with a capital efficient business strategy



\*Includes a single oncology indication from our Roche and Genentech collaboration.

and partnership value-drivers



### Data that is relatable and scalable is the Recursion differentiator

Recursion Data Universe: >21 PB of proprietary biological and chemical data, spanning phenomics, transcriptomics, invivomics, and more

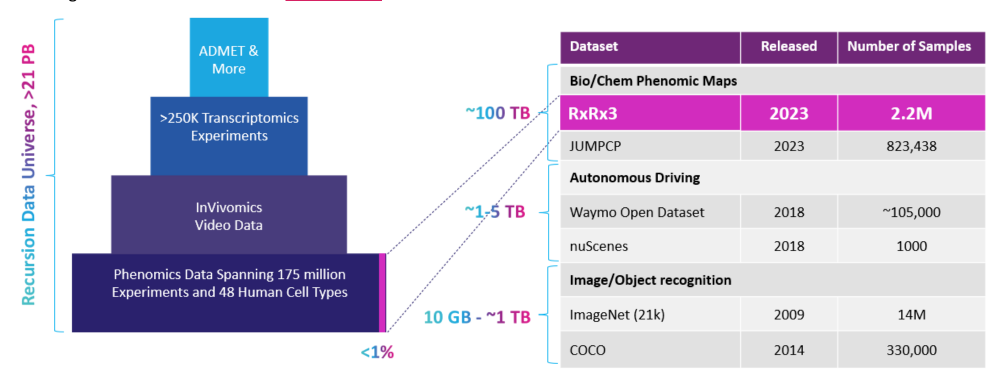
• We believe one of the largest biological and chemical datasets fit for the purpose of training large-scale ML models

**RXRX3:** CRISPR knockouts of most of the human genome, 1,600 FDA approved / commercially available bioactive compounds

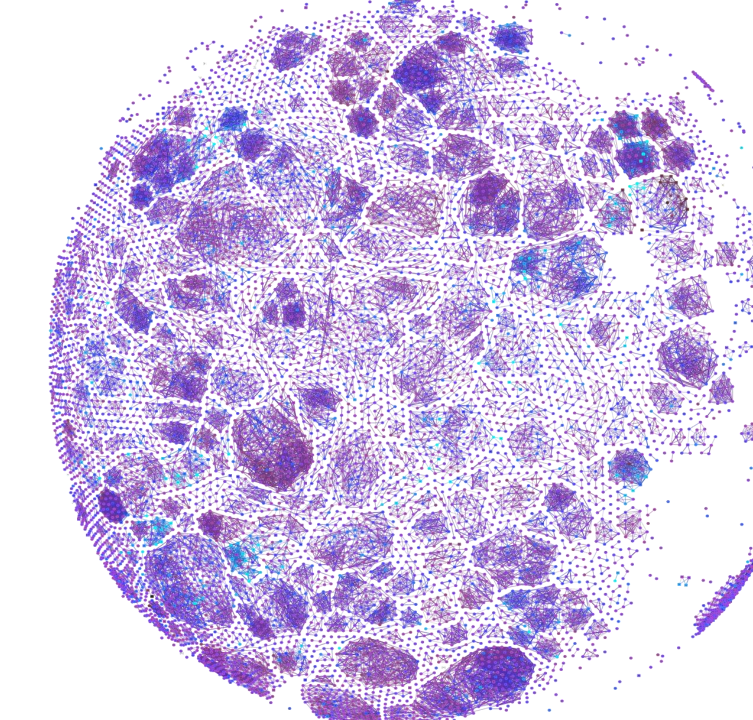
• We believe the largest public dataset of its kind, <1% of Recursion Data Universe, what Recursion can generate in ~1 week

MolRec™: freemium web-based application to explore compound and gene relationships in RXRX3

Start working with RXRX3 and MolRec™: www.rxrx.ai





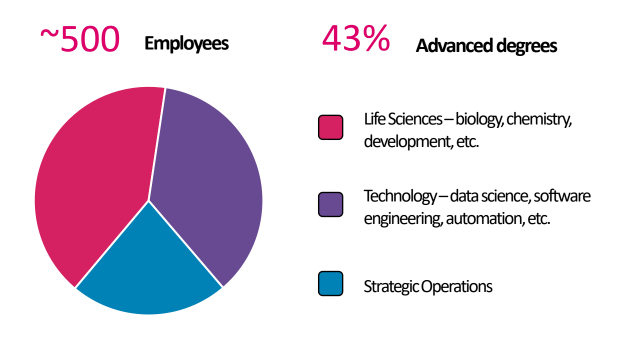






## What it takes to make this happen – a new kind of team and culture

#### **Team Members**



43% 55% 1% Female Male Non-Binary

Parity Pledge Signer gender parity and people of color parity

#### **ESG Highlights**

- ✓ Inaugural ESG report in 2022 reporting on Healthcare and Technology Metrics
- ✓ 100% of electricity powering our Biohive-1 supercomputer comes from renewable sources

#### **Community Impact**

altitude \_ lab

Founding Partner, Life Science Accelerator



Founding Member, Life Science Collective

#### **Committed to ESG Excellence**







### What to watch for at Recursion

#### **Upcoming Potential Milestones**

#### **Near-Term**

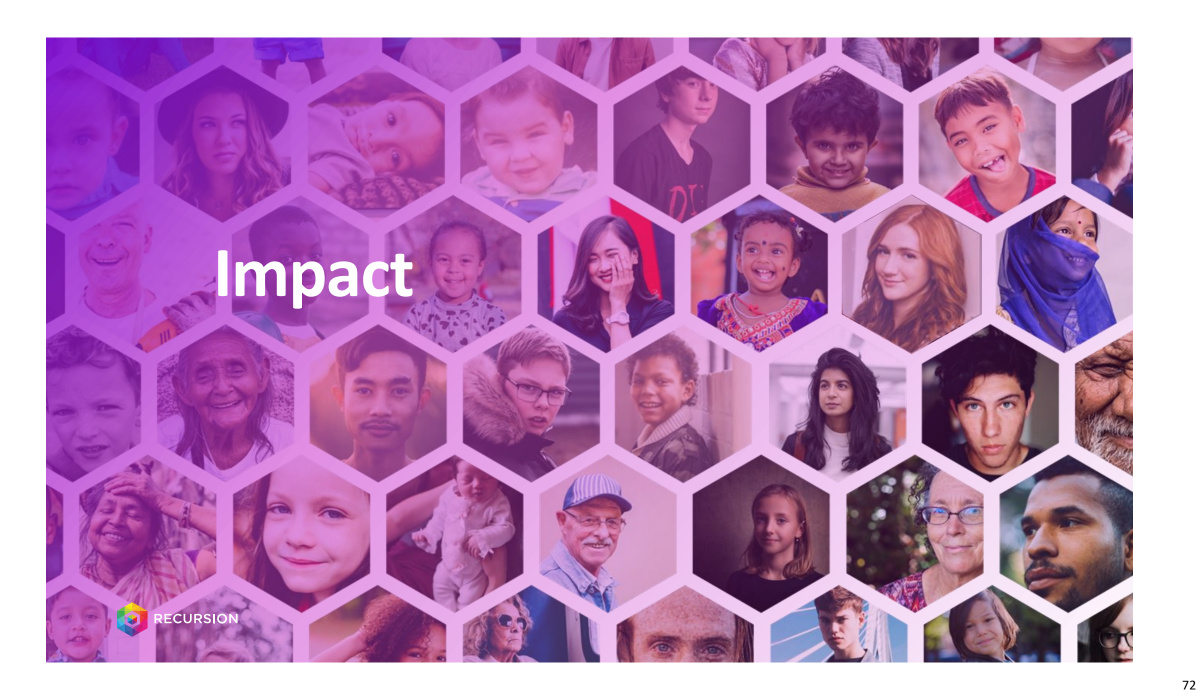
**Strong Financials** 

- Potential **option exercises** for partnership **programs**
- Potential option exercises for map building initiatives or data sharing
- Potential for additional partnership(s) in large, intractable areas of biology and / or technological innovation
- Ph1 clinical trial readout for C. difficile Infection program expected 2H 2023
- Potential for additional INDs and clinical starts, including
   Ph1b/2 trial initiation for AXIN1/APC program
- Potential for consolidation of technologies, talent and assets to accelerate the Recursion OS

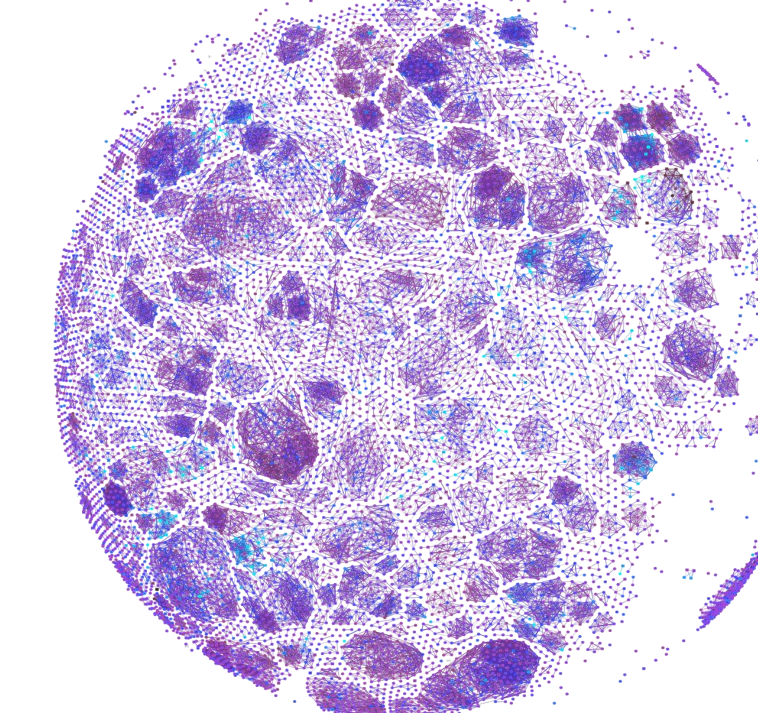
#### **Medium-Term**

- Multiple **POC readout(s)** for Al-discovered programs
  - NF2 interim safety analysis expected 2024
  - CCM top-line data expected 2H 2024
- Potential for additional INDs and clinical starts
- Potential option exercises for partnership programs
- Potential option exercises for map building initiatives or data sharing
- Potential additional partnership(s) in large, intractable areas of biology and / or technological innovation
- Recursion OS moves towards autonomous map building and navigation with digital and micro-synthetic chemistry

Learn more about Recursion's value proposition: <a href="https://www.recursion.com/download-day">www.recursion.com/download-day</a>



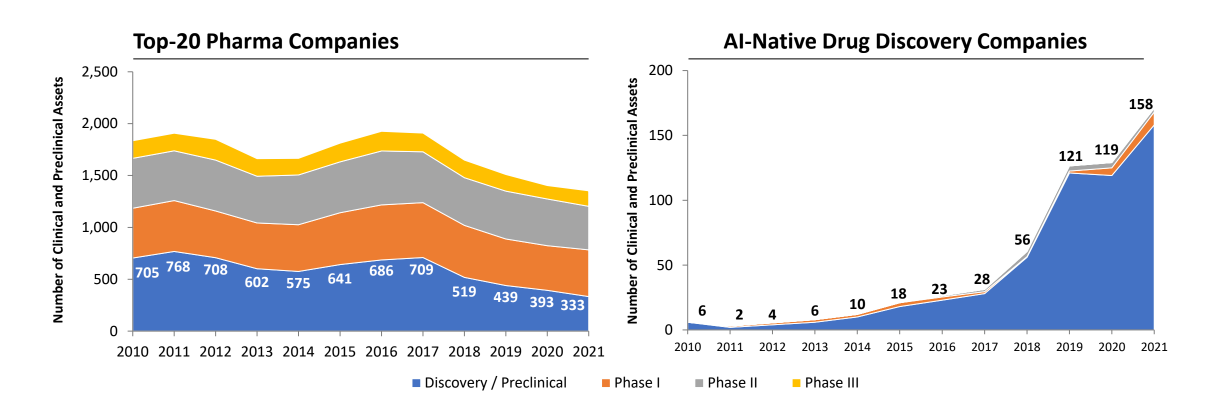
# Additional scientific and business context







# The biopharmaceutical industry faces pressure amidst declining efficiency in drug discovery

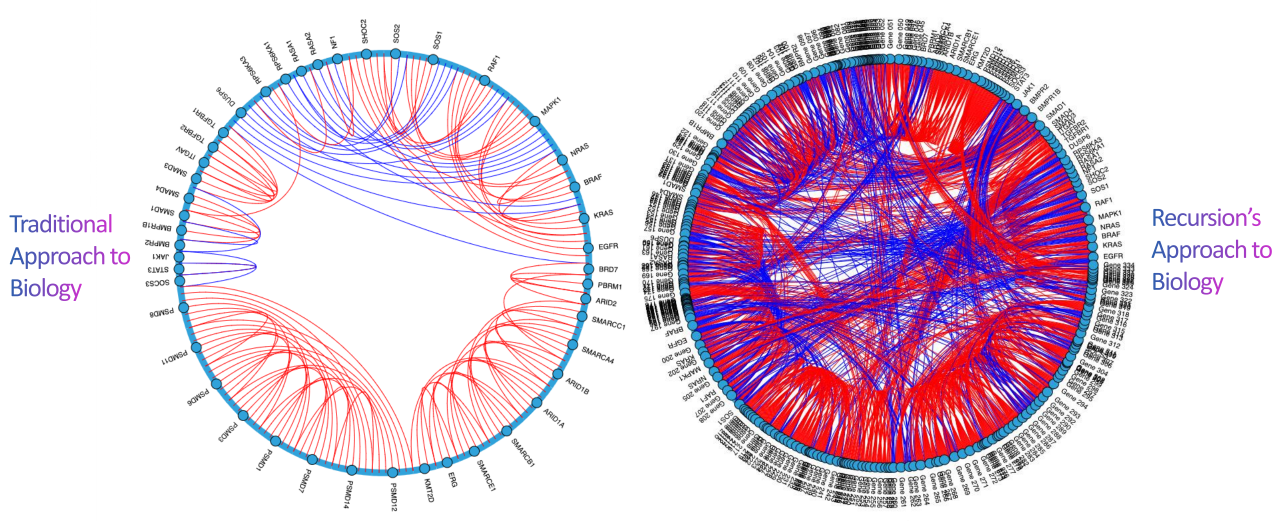


Al-enabled drug discovery efforts have proliferated alongside the declining efficiency of traditional approaches

Images adapted from Jayatunga, M., et al. Nature Reviews Drug Discovery 2022.



# Historical tools and the limits of human cognition have led to oversimplifying complex biological systems



#### Well-known primary relationships

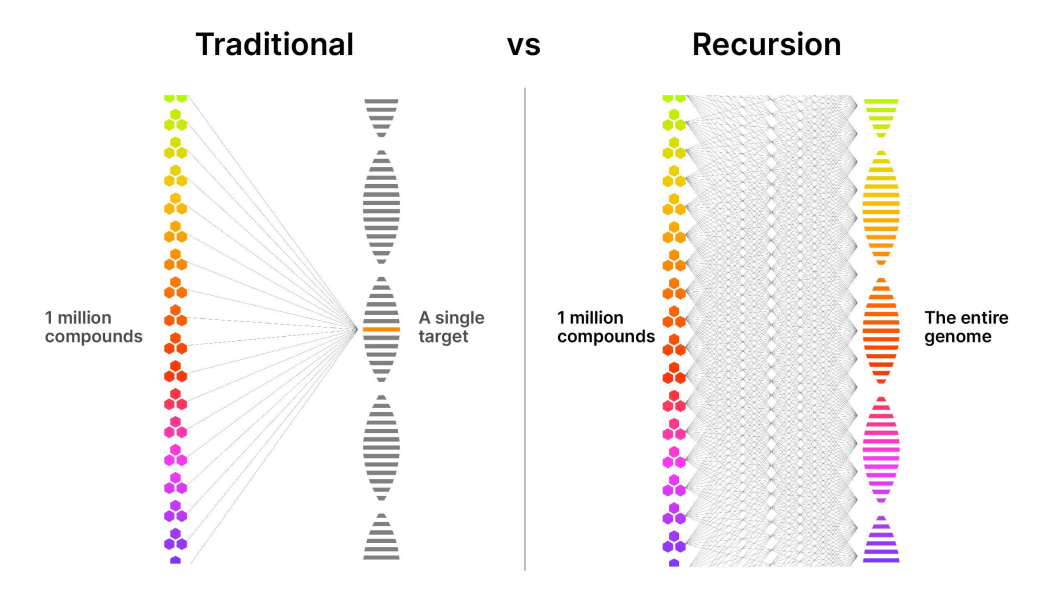
between key members of five pathways: JAK/STAT | SWI/SNF | TGFβ | RAS | Proteasome All primary relationships found by the Recursion OS

between key members of five pathways:

JAK/STAT | SWI/SNF | TGFβ | RAS | Proteasome

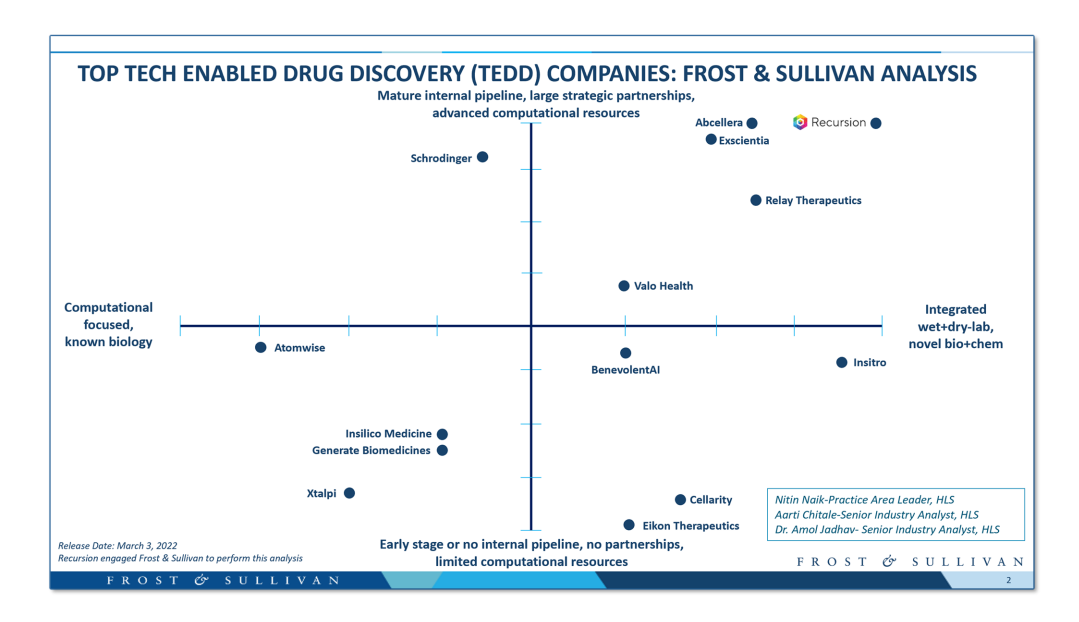


# Historical tools and the limits of human cognition have led to oversimplifying complex biological systems





### **Recursion is a leading TechBio company**





# Biology and chemistry are complex – data that is reliable, relatable, and scalable is the Recursion differentiator

Year	2018	2019	2020	2021	2022
Total Phenomics Experiments (Millions)	8	24	56	115	175
Total Transcriptomics Experiments (Thousands)	NA	NA	2	91	258
Data (PB)	1.8	4.3	6.8	12.9	21.2
Cell Types	12	25	36	38	48
Unique Compounds Physically Housed at Recursion <sup>1</sup> (Millions)	0.02	0.1	0.7	1.0	1.7
In Silico Chemistry Library (Billions)	NA	0.02	3	12	>1,000
Predicted Biological and Chemical Relationships <sup>2</sup> (Trillions)	NA	NA	0.01	0.2	3.1

<sup>&</sup>lt;sup>1</sup> Includes approximately 500,000 compounds from Bayer's proprietary library.

<sup>&</sup>lt;sup>2</sup> 'Predicted Relationships' refers to the number of Unique Perturbations that have been predicted using our maps.

### **COVID-19** research

Drug	Prediction	Correct?
Hydroxychloroquine	х	✓
Lopinavir	х	<b>✓</b>
Ritonavir	Х	<b>✓</b>
Remdesivir	<b>✓</b>	<b>✓</b>
Baricitinib	<b>✓</b>	<b>✓</b>
Tofacitinib	<b>✓</b>	<b>✓</b>
Ivermectin	Х	<b>✓</b>
Fluvoxamine	Х	<b>✓</b>
Dexamethasone	Х	X

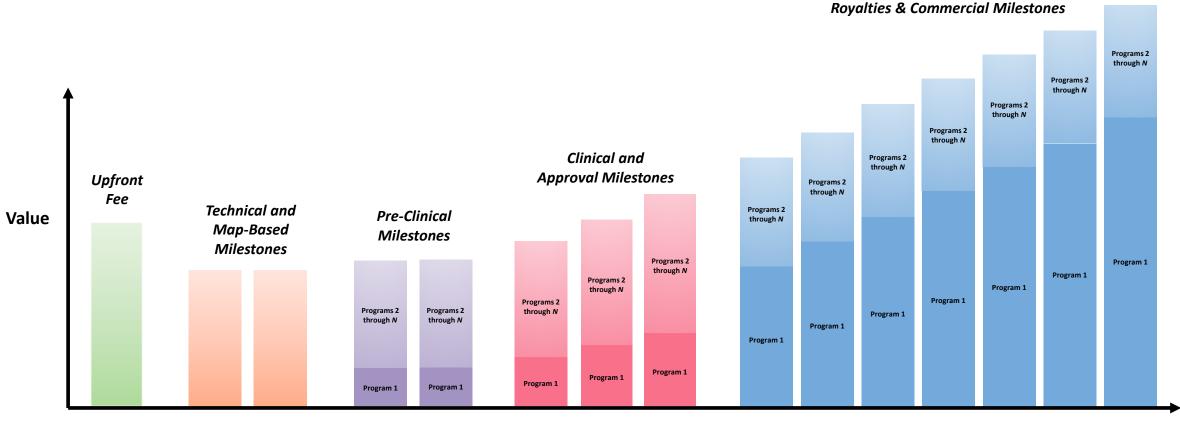
- Recursion conducted several AI-enabled experiments in April 2020 to investigate therapeutic potential for COVID-19
  - Included FDA-approved drugs, EMA-approved drugs, and compounds in late-stage clinical trials for the modulation of the effect of SARS-CoV-2 on human cells
- Experiments were compiled into the RxRx19 dataset
   (860+ GB of data) and made publicly available to
   accelerate the development of methods and pandemic
   treatments.
- Recursion OS correctly predicted 8 of 9 clinical trials associated with early and late-stage COVID-19

https://www.biorxiv.org/content/10.1101/2020.04.21.054387v1



# Transformational collaborations provide multiple potential value inflection points

Illustrative example of potential value inflection points



**Collaboration Timeline** 



# SYCAMORE Clinical Trial: REC-994 for CCM Phase 2 Underway

#### PREVALENCE & STANDARD OF CARE

~360,000

Symptomatic US + EU5, >1 million patients worldwide live with these lesions today

>5x larger US patient population than other rare diseases like Cystic Fibrosis (>31k patients)

#### No approved therapy

- No other potential therapeutic in industry-sponsored clinical development
- Most patients receive no treatment or only symptomatic therapy

#### CAUSE

LOF mutations in genes CCM1, CCM2 & CCM3, key for maintaining the structural integrity of the vasculature due to unknown mechanisms

#### PATHOPHYSIOLOGY & REASON TO BELIEVE

Vascular malformations of the CNS leading to focal neurological deficits, hemorrhage and other symptoms



Efficacy in Recursion OS as well as functional validation via scavenging of massive superoxide accumulation in cellular models; reduction in lesion number with chronic administration in mice

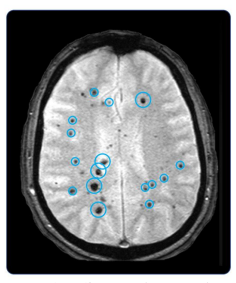






- Targeting sporadic and familial symptomatic CCM patients with CCM1, CCM2, and CCM3 mutations
- Superoxide scavenger, small molecule

- Phase 2 trial initiated in Q1 2022
- US & EU Orphan Drug Designation
- Oral dosing



Vascular malformations (cavernomas)



Julia - living with CCM



## POPLAR Clinical Trial: REC-2282 for NF2 Phase 2/3 Underway

**PREVALENCE & STANDARD OF CARE** 

~33,000

Treatable US + EU

#### No approved therapy

- There are no approved drugs for NF2
- Surgery is standard of care (when feasible)
- Location may make complete resection untenable, leading to hearing loss, facial paralysis, poor balance and visual difficulty

#### **CAUSE**

**LOF mutations in NF2 tumor suppressor gene**, leading to deficiencies in the tumor suppressor protein merlin

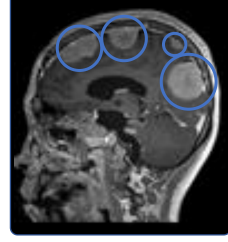
#### PATHOPHYSIOLOGY & REASON TO BELIEVE

Inherited rare **CNS tumor syndrome** leading to loss of hearing and mobility, other focal neurologic deficits



Efficacy in Recursion OS, cellular, and animal models; suppression of aberrant ERK, AKT, and S6 pathway activation in a Phase 1 PD Study in NF2 patient tumors





Intracranial meningiomas



**KEY ELEMENTS** 

- Targeting familial and sporadic NF2 meningioma patients
- HDAC inhibitor, small molecule
- Oral dosing

- Phase 2/3 trial initiated in Q2 2022
- Fast-Track and US & EU Orphan Drug Designation

Ricki - living with NF2



### **TUPELO Clinical Trial: REC-4881 for FAP Phase 2 Underway**

**PREVALENCE & STANDARD OF CARE** 

~50,000

Diagnosed US + EU5

#### No approved therapy

**KEY ELEMENTS** 

- Colectomy during adolescence (with or without removal of rectum) is standard of care
- Post-colectomy, patients still at significant risk of polyps progressing to GI cancer
- Significant decrease in quality-of-life post-colectomy (continued endoscopies, surgical intervention)

#### **CAUSE**

Inactivating mutations in the tumor suppressor gene APC

#### PATHOPHYSIOLOGY & REASON TO BELIEVE

**Polyps throughout the GI tract** with extremely high risk of malignant transformation



Efficacy in the Recursion OS showed specific MEK 1/2 inhibitors had an effect in context of *APC* LOF. Subsequent APC<sup>min</sup> mouse model showed potent reduction in polyps and dysplastic adenomas



- MEK inhibitor, small molecule
- Oral dosing

- Phase 2 trial initiated in Q3 2022
- Fast-Track and US & EU Orphan Drug Designation



Polyps Found in Colon and Upper GI Tract

### **Clinical Program: REC-4881 for AXIN1 or APC Mutant Cancers**

**PREVALENCE & STANDARD OF CARE** 

~65,000

Treatable US + EU5

**Substantial need** for developing therapeutics for patients harboring mutations in *AXIN1* or *APC*, as **these mutations are considered undruggable** 

To our knowledge, REC-4881 is the **only industry sponsored small molecule therapeutic** designed to enroll solid tumor patients harboring mutations in *AXIN1* or *APC* 

**CAUSE** 

LOF mutations in AXIN1 or APC tumor suppressor genes

#### **PATHOPHYSIOLOGY & REASON TO BELIEVE**

Alterations in the WNT pathway are found in a wide variety of tumors and confer poor prognosis and resistance to standard of care



Efficacy in the Recursion OS and favorable results in PDX models harboring *AXIN1* or *APC* mutations vs wild-type leading to a significant PFS benefit in HCC and Ovarian tumors





- Targeting solid tumors with AXIN1 or APC mutant cancers
- MEK inhibitor, small molecule
- Oral dosing

- Finalize design of a Phase 1b/2 biomarker-enriched trial
- Initiate Phase 1b/2 trial in select tumor types in early 2024



Gross morphology of HCC



#### Clinical: C. Difficile

### Clinical Trial: REC-3964 for C. Difficile Phase 1 Underway

**PREVALENCE & STANDARD OF CARE** 

~730,000

Diagnosed US + EU5

Standard of care includes antibiotic therapies which can further impair gut flora, and lead to relapse

#### **CAUSE**

**C.** difficile toxins from colonizing bacterium causes degradation of colon cell junction, toxin transit to bloodstream, and morbidity to host

#### **KEY ELEMENTS**

- Selective C. diff toxin inhibitor, small molecule
- Non-antibiotic approach with potential for combination with SOC and other therapies
- Designed for selective antitoxin pharmacology to target infection
- FIH Phase 1 trial initiated in Q3 2022

#### PATHOPHYSIOLOGY & REASON TO BELIEVE

Highly recurrent infectious disease with severe diarrhea, colitis, and risk of toxic megacolon, sepsis, and death



Recursion OS identified a new chemical entity for recurrent C. difficile infection and potentially prophylaxis via glycosyl transferase inhibition with potential to be orally active



#### TRIAL UPDATE

- Enrollment is progressing
- In several SAD cohorts and at least one MD cohort, REC-3964 has been extremely safe and well tolerated
- Complete safety and PK data readout expected 2H 2023



Colleen - lived with rCDI