



BIO Digital June 2021

Forward-Looking Statements

This presentation contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of 1995. Any statement in this document that is not a historical fact is a "forward-looking statements" within the meaning of Section 27A of the Securities Act and Section 21E of the Securities Exchange Act and are usually identified by the use of words such as "anticipates," "believes," "estimates," "expects," "intends," "may," "plans," "projects," "seeks," "should," "will," and variations of such words or similar expressions. We intend these forward-looking statements to be covered by the safe harbor provisions for forward-looking statements contained in Section 27A of the Securities Act and Section 21E of the Securities Exchange Act of 1934, as amended, and are making this statement for purposes of complying with those safe harbor provisions. These forward-looking statements reflect our current views about our plans, intentions, expectations, strategies and prospects, which are based on the information currently available to us and on assumptions we have made. Although we believe that our plans, intentions, expectations, strategies and prospects as reflected in or suggested by those forward-looking statements are reasonable, we can give no assurance that the plans, intentions, expectations or strategies will be attained or achieved. Furthermore, actual results may differ materially from those described in the forward-looking statements and will be affected by a variety of risks and factors that are beyond our control.

Risks and uncertainties for Plus include, but are not limited to: an inability or delay in obtaining required regulatory approvals for product candidates, which may result in unexpected cost expenditures; risks inherent in drug development in general; uncertainties in obtaining successful clinical results for product candidates and unexpected costs that may result therefrom; failure to realize any value of certain product candidates developed and being developed in light of inherent risks and difficulties involved in successfully bringing product candidates to market; inability to develop new product candidates and support existing products; the approval by the FDA and any other similar foreign regulatory authorities of other competing or superior products brought to market; risks resulting from unforeseen side effects; risk that the market for the combined company's products may not be as large as expected; inability to obtain, maintain and enforce patents and other intellectual property rights or the unexpected costs associated with such enforcement or litigation; inability to obtain and maintain commercial manufacturing arrangements with third-party manufacturers or establish commercial scale manufacturing capabilities; loss of or diminished demand from one or more key customers or distributors; unexpected cost increases and pricing pressures; economic recession and its negative impact on customers, vendors or suppliers; uncertainties of cash flows, expenses and inability to meet working capital needs; and other risks and uncertainties detailed in the risk factors section of Plus' Form 10-K and Forms 10-Q filed with the SEC, as well as other filings Plus makes with the SEC from time-to-time. Many of these factors that will determine actual results are beyond Plus' ability to control or predict. Plus disclaims any obligation to update information contained in these forward-looking statements, whether as a result of new information, future events or otherwise, except as required by law.



We believe in the critical importance and urgency in developing and delivering complex, innovative treatments for patients battling rare and CNS cancers.



CNS & Rare Cancers

Responsible for Substantial Morbidity and Mortality Worldwide

- + Significant unmet medical needs
 - + 5-year survival of 36%
 - + Few approved treatments
- + Rare diseases
 - + FDA ODD eligible
 - + Sizeable aggregate population
- + Biological overlap



This accounts for **70,000 to 170,000** new patients diagnosed with CNS metastases each year.

FACTS ABOUT BRAIN TUMORS

An estimated **700,000+ PEOPLE** in the U.S. are living with a primary brain or central nervous system (CNS) tumor diagnosis:

210,000 WITH MALIGNANT TUMORS 490,000 WITH NON-MALIGNANT TUMORS

APPROXIMATELY

6-28%

OF ALL OTHER CANCERS LATER
DEVELOP A CNS METASTASES



NON-MALIGNANT TUMORS (58,860 Cases)

IN 2020, NEW PRIMARY BRAIN TUMOR DIAGNOSES INCLUDED



70% 30%

MALIGNANT TUMORS (24,970 CASES)

PRIMARY BRAIN TUMOR TYPES



- Meningioma (38%)
- Pituitary (17%)
- Glioblastoma (15%)
- Other Non-Glioma (11%)
- Other Glioma (11%)
- Nerve Sheath (9%)

4

Sources: CBTRUS Statistical Report, NBTS

Glioblastoma

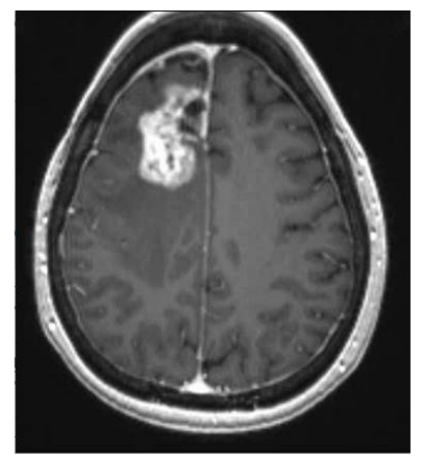
A Rare, Incurable, and Fatal Brain Cancer with No Good Treatment Options



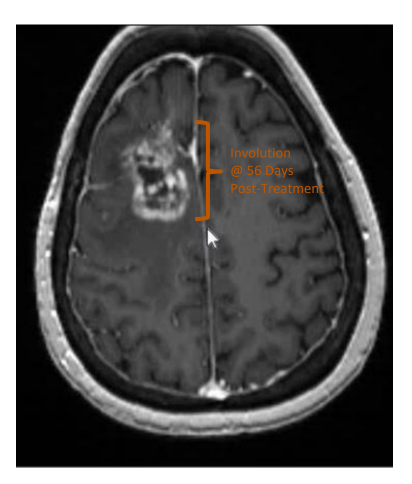


PLUS Therapeutics: Encouraging Phase I Data

Patient with Recurrent Glioblastoma- Before and After Tumor Ablation Following High Dose RNL Therapy



Before Treatment



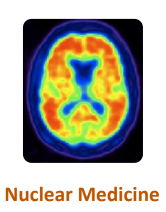
56 Days Post-Treatment



PLUS Therapeutics: A Novel Approach to Intracranial Neoplasms

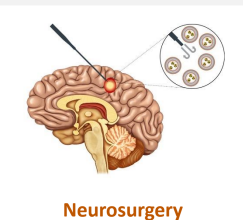
Marriage of New Developments Across Multiple Specialities







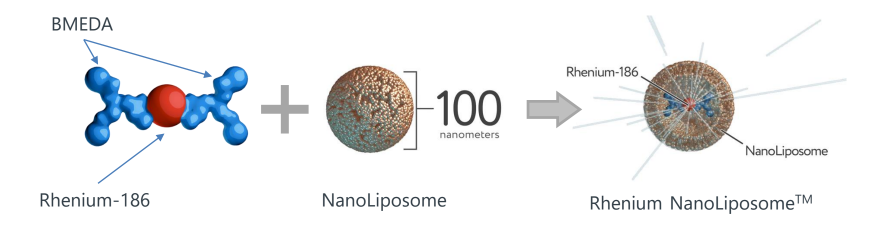






Lead Asset: Rhenium Nanoliposome or RNL™

Proprietary Nanoscale Compound with a Unique Isotope



RHENIUM 186

- Dual energy emitter- Beta (cytotoxic) & Gamma (imaging)
- Short average path length- precision
- Low dose rate- safer for normal tissues
- High radiation density- overwhelms innate DNA repair mechanisms



PLUS Therapeutics: Pipeline

Growing Preclinical and Clinical Therapeutic Pipeline

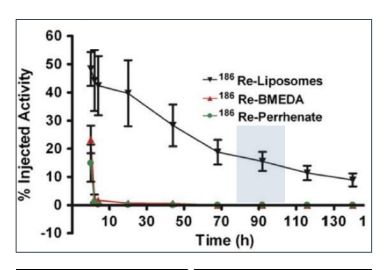
	Investigational Drug	Indication	Designation	Stage	External Funding
		Recurrent Glioblastoma	FDA Orphan Drug FDA Fast Track	Phase 1	NIH/NCI to PII
• •	Rhenium NanoLiposome (RNL™)	Leptomeningeal Cancer	-	IND pending	-
		Pediatric CNS Cancers: Ependymoma, High Grade Glioma & Diffuse Intrinsic Pontine Glioma	-	IND pending	-
Combination Therapy: Chemotherapeutic & Radiotherapeutic	Doxorubicin & Rhenium NanoLiposome (DRNL™)	Multiple	-	Preclinical	-

Plus DocePLUS[™] and DoxoPLUS[™] clinical-stage chemotherapeutic assets available for partnering



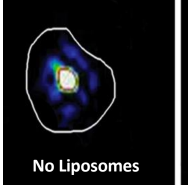
Spatiotemporal Behavior of RNL™ Following Brain Delivery

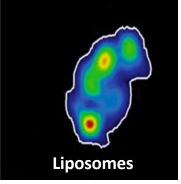
Prolonged Half-Life and Brain Retention



Tumor Retention

Liposomal encapsulation significantly extends the *in vivo* intracranial half-life of rhenium-186 (90 hours) and decreases clearance rate from the brain.





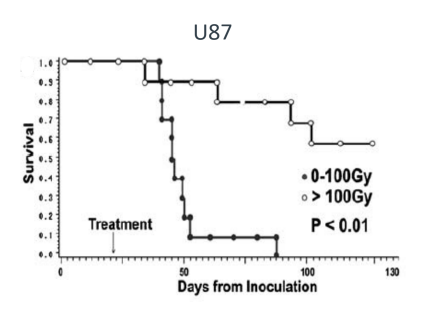
Tumor Dispersion

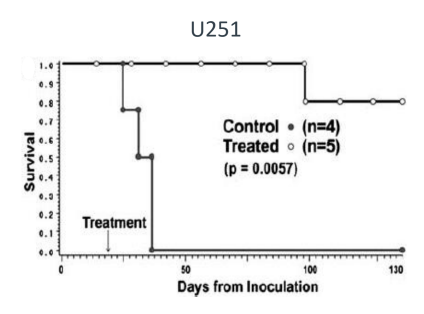
Liposomal encapsulation significantly extends rhenium-186 retention within the tumor and therefore improves dispersion characteristics in tissues.



RNL™ Preclinical Data

RNL Significantly Prolongs Survival in U87 & U251 Intracranial Xenograft Models





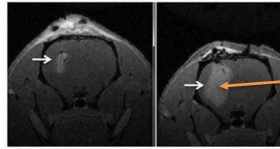
- Doses of up to 1,845 Gy were tolerated without weight loss or neurological deficit.
- No maximum tolerated dose of RNL reached.
- Statistically significant prolongation in survival, limited only be the end of the experiment.
- Blinded histologic analysis by neuropathologist showed no residual tumor all treated animals.



RNL™ Preclinical Data

Tumor Regression in U87 & U251 Intracranial Xenograft Models

Control



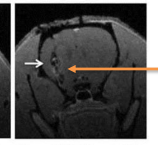
Day 14 Control

& expansion in untreated control animals with brain cancer.

Tumor growth

¹⁸⁶Re-Liposome Treatment





Tumor regression in RNL™ treated brain cancers.

Day -1

Day 14 Post-Tx Day 28 Post-Tx Day 70 Post-Tx

- Bioluminescence assay showed many of the treated animals had a loss of activity to background levels suggesting complete eradication of the tumor.
- MRI analysis (above) supported the observation of tumor eradication.
- Blinded histologic evaluation by neuropathologist showed no residual disease.



Day -1

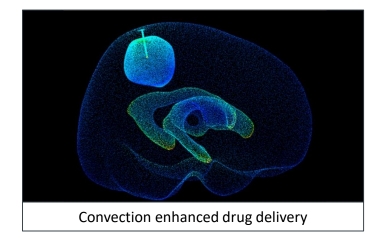
RNL™ Fits Within Modern Clinical Workflows

Plan, Administer, and Confirm RNL™ Treatment with the Latest Imaging & Surgical Techniques











RNL™ for Recurrent Glioblastoma

Potential Advantages Compared to External Beam Radiation Therapy







RNL™ for Recurrent Glioblastoma

Phase 1/2 Clinical Trial Design

Multi-center, sequential cohort, open-label, volume and dose finding study of the safety, tolerability, and distribution of ¹⁸⁶RNL given by convection enhanced delivery to patients with recurrent or progressive malignant glioma after standard surgical, radiation, and/or chemotherapy treatment.

- + Single arm, prospective Phase 1/2 study utilizing a modified Fibonacci dose escalation scheme, followed by an expansion at the designated recommended phase 2 dose (RP2D).
- + Maximum number of planned subjects: up to 55 subjects (including patients enrolled in the Phase 1 dose escalation trial and a subsequent cohort at the RP2D).
- + Supported by a NIH/NCI grant through Phase 2.













Phase 1 Dosing Scheme

Cohort	Infused Volume (mL)	Total RNL™ Activity (mCi)	Concentration (mCi/mL)	Average Absorbed Dose (Gy)	Status
1	0.66	1.0	1.5	198	
2	1.32	2.0	1.5	122	
3	2.64	4.0	1.5	234	First 7 cohorts
4	5.28	8.0	1.5	171	complete
5	5.28	13.4	2.5	423	(n = 21 subjects)
6	8.80	22.3	2.5	287	
7	8.80	22.3	2.5	584	

^{*}Cohort 7 utilized same volume and dose as cohort 6 but with increase in maximum flow rate to 20 microliters/minute

Tumor Coverage Implication:

+ 8.8 mL treatment volume may cover ~3.5 cm diameter tumor.





Safety Summary

RNL Is Safe and Well Tolerated

Thus far, in the Phase I study of 21 subjects with recurrent Glioblastoma (GBM) receiving a single dose of RNL™:

- RNL™ appears to be safe and well-tolerated.
- There have been no dose limiting toxicities.
- The majority of AEs reported were mild or moderate (Grade 1 or 2) in intensity.
- Most AEs were considered causally unrelated to RNL™ except scalp discomfort, which was considered related to the surgical procedure.
- No meaningful differences or patterns in the incidence of related treatment emergent adverse effects reported across individual treatment groups or cohorts.
- Neither the incidence nor severity of AEs appeared to increase with increasing doses of RNL™.



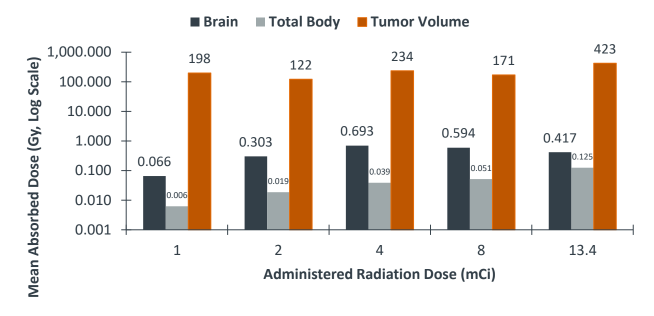


Dosimetry Evaluation

High Absorbed Doses to Tumor Volume, Not Brain or Total Body

- + Absorbed Dose (Gy) to Tumor Volume
 - + Mean 239 Gy (range 9-593 Gy)
- + Ratio of mean tumor volume AD/mean total body AD in cohorts 4 & 5, ≥ 3,000

Mean Absorbed Dose To Brain, Total Body, and Tumor Volume







Clinical Data Update

Continued Positive Delivery & Efficacy Signals Through Cohort 7

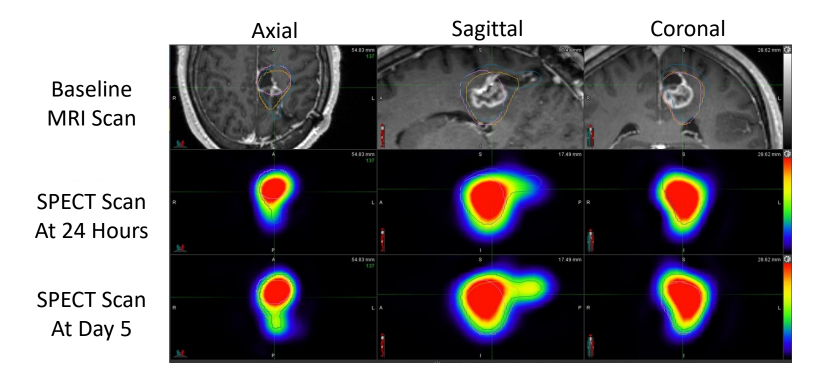
- + Successfully deliver up to 740 Gray of radiation to tumors
- + Patients receiving prior Bevacizumab (n = 5) poorly convect the RNL through the tumor
- + Average tumor coverage is 77% in 16 Bevacizumab naïve patients and increasing with volume & dose escalation
- + Overall survival results continue to trend positively
- + Thirty-three percent or 7/21 patients remain alive





Tumor Coverage and Retention

Subject 01-014



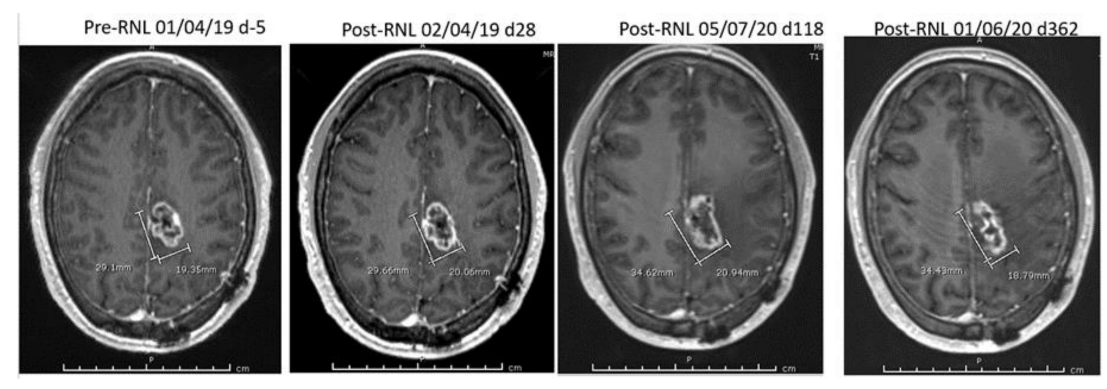
- + Tumor volume was 6.5 mL and tumor coverage was > 90%
- + Absorbed dose delivered to tumor was 419 Gy





Natural History of Recurrent GBM Lesions After RNL™

Subject 01-014: Tumor Response Observed to Day 362



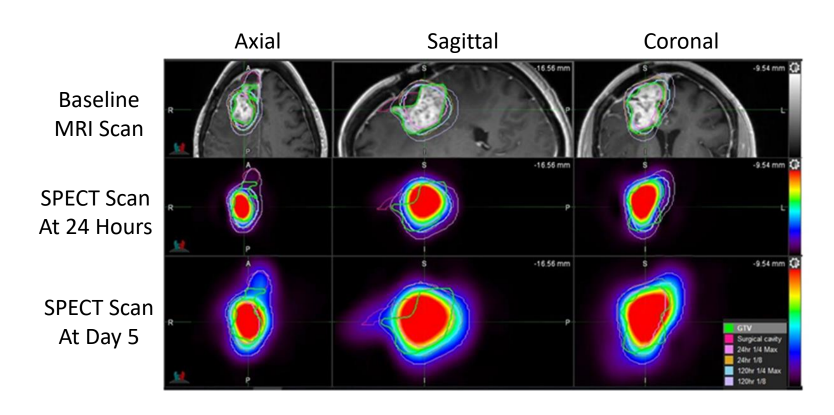
- + MRI scans revealed an initial increase in size which peaked at Day 118, with some associated edema, followed by tumor shrinkage out to at least Day 362
- + Patient remains alive (>600 days)





Tumor Coverage and Retention

Subject 01-017



- + Tumor volume was 18.8 mL and tumor coverage was 87%
- + Absorbed dose delivered to tumor was 336 Gy

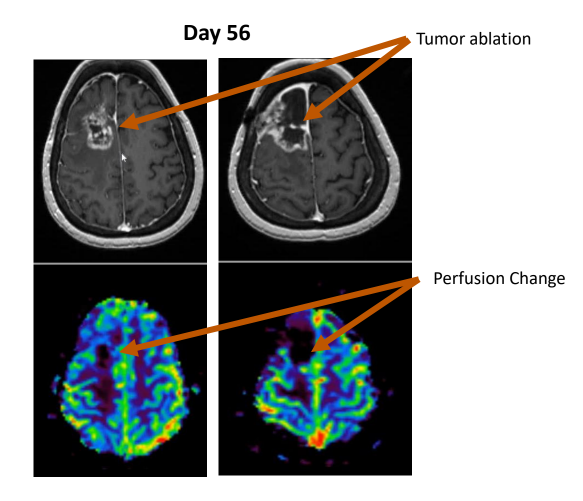




MRI & Perfusion Imaging

Subject 01-017: Tumor Ablation & Loss of Perfusion

Pre-Treatment





RNL™ Indication Expansion

Opportunity to Address Significant Unmet Needs for Multiple Rare Cancers

Leptomeningeal Carcinomatosis (LMC)

+ U.S. annual incidence: 110,000

+ Derived from breast/lung cancers, melanoma, GI malignancies

+ No clear SoC: radiation & chemo used

+ 5-year survival: unreported

+ Survival if untreated: 4-6 weeks

+ Survival if treated: 4-6 months

Peritoneal Carcinomatosis

+ U.S. annual incidence: 72,000

+ Derived from colorectal/gastric/ovarian/pancreatic cancers

+ Controversial SoC: chemo & cytoreduction surgery

+ 5=year survival <3%-66%

Pediatric Brain Cancer

+ U.S. annual incidence: 1,000*

+ Current SOC: steroids, radiation, surgery &/or chemo

+ 5-year survival: <3% (DIPG), <20% (HGG), 75% (Ependymoma)

+ Median survival: 10-11 months (DIPG)

Head & Neck Squamous Cell Carcinoma

+ U.S. annual incidence: 53,000

+ Current SoC (locally advanced tumors): radiation, surgery, systemic therapy

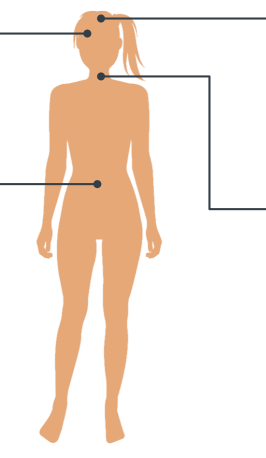
+ Current SoC (small tumors without nodal involvement): radiation or surgery

+ 5-year survival: 66%

+ 30-40% develop recurrent locoregional cancer

+ 20-30% develop metastatic disease

+ Median OS in recurrent or metastatic: 6-15 months





^{*} High Grade Glioma, Ependymoma, Diffuse Intrinsic Pontine Glioma

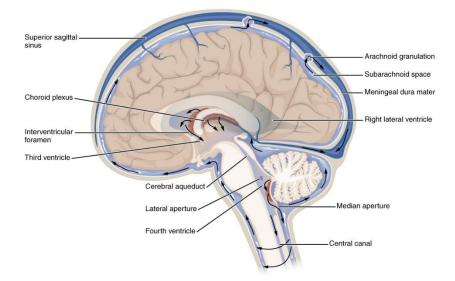
RNL™ Rationale for Leptomeningeal Cancer

Disease Background

+ Leptomeningeal cancer, also known as carcinomatosis, is a cancer that starts in one part of the body spreads to the leptomeningeal lining of the brain and spinal cord surrounding the CSF space.

100 nm NanoLiposomes in CSF

- + Circulate feely throughout the CSF.
- + Migrate to meningeal surfaces where LMC is located.
- + Have an extended half life several weeks vs. hours with unencapsulated drugs.



Preclinical Data

+ Unpublished data suggests RNL may be effective in preclinical models of leptomeningeal cancer



Pre IND Meeting Feedback on Potential New RNL Indications

Leptomeningeal Disease

- Additional GLP toxicology studies are not needed to support initiation of the pediatric study.
- They provided constructive feedback on the study synopsis that should be helpful as a full protocol is developed.
- They provide helpful guidance on the data required for dose justification and suggest that we request a formal meeting to review division recommended phase 2 dose (RP2D) and prior to evaluation of expansion cohorts or trials designed to demonstrate efficacy and safety

Pediatric Brain Cancer Disease

- Additional GLP toxicology studies are not needed to support initiation of the pediatric study.
- They provided constructive feedback on the study synopsis that should be helpful as a full protocol is developed.
- They are requesting the study design be modified to include separate dose escalation components for each proposed anatomic tumor location.



RNL™ Development Plan

Recurrent Glioblastoma	2021	2022	2023	2024	2025
ReSPECT™ Phase 1 for rGBM					
Manufacturing & Scale Up for PIII					
Potential P II/Expansion arms	////				
Anticipated Phase III for rGBM					

LMC & Pediatric Brain Cancer	2021	2022	2023	2024	2025
FDA IND Filed LMC					
Pilot/Phase 1 for LMC					
FDA IND Meeting for Pediatric Brain Cancer					
Pilot/Phase 1 for Pediatric Brain Cancer					



Forthcoming Milestones for 2021 and into 2022

- Complete enrollment & report data from U.S. ReSPECT™ Phase I trial for RNL™ in recurrent GBM
- DSMB meeting and FDA Complete CMC meetings
- Complete CMC activities for RNL™ for a phase III drug supply
- Potentially proceed with Phase 2 and/or ongoing dose escalation
- Hove RNL™ into Phase I for leptomeningeal cancer 1st and & a pediatric brain cancer indication to follow
- **+** Complete additional preclinical studies for RNL™ and DRNL™
- Potential acquisition, in-license new drug development candidates
- Partner RNL™, DocePLUS™ & DoxoPLUS™ assets if possible



Capitalization Summary

Select Data (as of March 31, 2021)				
Cash	\$14.4M			
Common Shares Outstanding	10,180,525			
Series U warrants	2,141,000			
Senior Term Loan Principal (matures 2024)	\$4.3M			





+ Headquarters: Austin, Texas, USA

+ Manufacturing: San Antonio, Texas, USA

+ Nasdaq: PSTV

+ Corporate Website: PlusTherapeutics.com

+ ReSPECT™ Website: ReSPECT-Trials.com

