



### FORWARD-LOOKING STATEMENTS

Statements in this presentation contain "forward-looking statements" that are subject to substantial risks and uncertainties. Forward-looking statements contained in this presentation may be identified by the use of words such as "anticipate," "expect," "believe," "will," "may," "should," "estimate," "project," "outlook," "forecast" or other similar words, and include, without limitation, statements regarding Annovis Bio, Inc. 's expectations regarding projected timelines of clinical trials, and expectations regarding current or future clinical trials. Forward-looking statements are based on Annovis Bio, Inc.'s current expectations and are subject to inherent uncertainties, risks and assumptions that are difficult to predict. Further, certain forward-looking statements are based on assumptions as to future events that may not prove to be accurate, including that clinical trials may be delayed; that the data reported herein is interim data, conclusions as to which may be superseded by subsequent data we expect to receive in connection with Phase 2a trials and/or subsequent clinical trials; and that any anticipated meeting with or presentation to the FDA may be delayed. These and other risks and uncertainties are described more fully in the section fitled "Risk Factors" in the Annual Report on Form 10-K for the year ended December 31, 2020 and other reports filed with the Securities and Exchange Commission. Forward-looking statements contained in this presentation are made as of this date, and Annovis Bio, Inc. undertakes no duty to update such information except as required under applicable law.

### FREE WRITING PROSPECTUS

Annovis Bio, Inc. has filed a registration statement (including a prospectus) with the SEC for the offering to which this communication relates. Before you invest, you should read the prospectus in that registration statement and other documents Annovis has filed with the SEC for more complete information about Annovis and this offering. You may get these documents for free by visiting EDGAR on the SEC Web site at www.sec.gov. Alternatively, the issuer, any underwriter or any dealer participating in the offering will arrange to send you the prospectus if you contact ThinkEquity, a division of Fordham Financial Management, Inc., located at 17 State Street, 22nd Floor, New York, New York 10004, by telephone at (877) 436-3673, or by email at prospectus@think-equity.com.



## **HIGHLIGHTS**

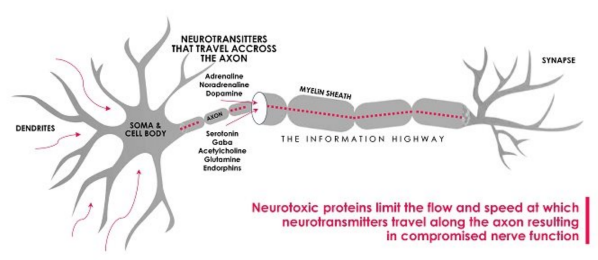
A novel approach to treat neurodegeneration is desperately needed

- Annovis is developing drugs for Alzheimer's (AD) and Parkinson's disease (PD), including the orphan indication Alzheimer's in Down Syndrome (AD-DS)
- Lead compound ANVS401 in Phase 2a clinical trial, is the only drug to improve cognition in AD and motor function in PD patients, as recently announced
- ANVS401 reduced inflammation, in PD patients as recently announced. Additional biomarker data to come.
- Successful completion of phase 2a clinical trials will validate our approach and allow start of two phase 3 studies



# HOW NERVE CELLS WORK

**In healthy nerve cells** little packages containing neurotransmitters or nerve growth factors travel unimpaired from the cell body through the axon to the synapse.





# NEUROTOXIC PROTEINS IMPAIR AXONAL TRANSPORT AND CAUSE A TOXIC CASCADE

# HIGH LEVELS OF NEUROTOXIC PROTEINS

IMPAIRED AXONAL TRANSPORT

SLOWER SYNAPTIC TRANSMISSION

INFLAMMATION

**DEATH OF NERVE CELLS** 

LOSS OF COGNITIVE AND MOTOR FUNCTION

# ANVS401 LOWERS LEVELS OF NEUROTOXIC PROTEINS

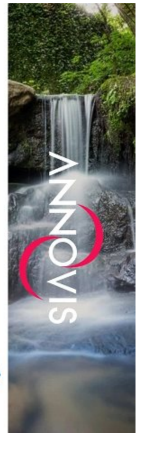
IMPROVED AXONAL TRANSPORT

INCREASED SYNAPTIC TRANSMISSION

NO INFLAMMATION

**HEALTHY NERVE CELLS** 

IMPROVED COGNITIVE AND MOTOR FUNCTION



ANVS401 IMPROVES AXONAL TRANSPORT AND IMPEDES THE TOXIC CASCADE

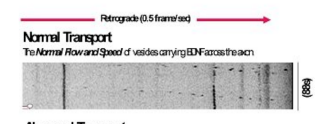


### NEURODEGENERATION IS AN AXONAL TRANSPORT DISEASE

"Axonal transport disruption is linked to human neurological conditions." - Nature Review, September 2019

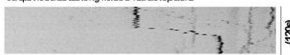
### Axonal transport is responsible for:

- Neurotransmitters GABA (anxiety), ACh (cognition), dopamine (movement), serotonin (mood)
- Neurotrophic factors NGF, BDNF
- All communication within and between nerve cells



Abnormal Transport

Stockte Bokage and Soving of EDF arcostreach Back area demonstrate where transport is sloved due to high levels of neurotoic podeins



TREATED WITH ANV\$ 401
Tre Flow and Speed of avoral transport is improved.

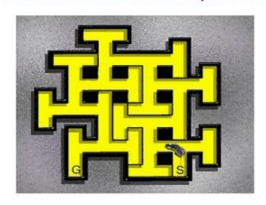


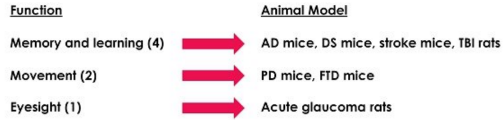
APP, Ab-2, CSB— Mittley, UCSD (SBN—Issacon, Hirvard, Lee; URmn, Tau— UMLerich & Zerich, Ht.— Mittley, UCSD, TDRS— Taylor, Northwestern



# RESULTS IN ANIMALS

Multiple animal studies showed that ANVS401 improved the affected function







# TWO PHASE 2 CLINICAL TRIALS

	AD Trial	PD Trial	
CRO	Parexel		
Therapeutic Area	Early to Moderate AD	Early to Moderate PD	
Phase	2	2	
Patients	14	14 + 40	
Sites	12	12	
Country	United States		
Design	Double-Blind, Placebo-Controlled, Biomarker Study		
Endpoints	Reversal of Toxic Cascade		



# PHASE 2 CLINICAL TRIAL IN AD AND PD TO MEASURE REVERSAL OF THE TOXIC CASCADE AND IMPROVEMENT IN BRAIN FUNCTION

### **ENDPOINTS**

TARGET: DECREASE IN NEUROTOXIC PROTEINS

PATHWAY: INCREASE IN NEUROTRANSMITTERS

LOWERING OF INFLAMMATORY PROTEINS

LOWERING OF NEURODEGENERATION

MARKERS

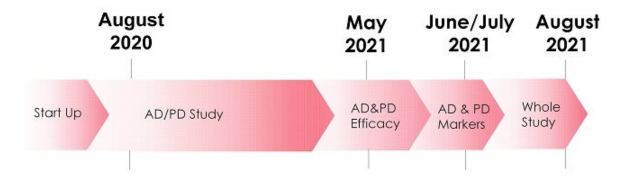
**EFFICACY:** COGNITION AND MOTOR FUNCTION





# TIMELINE OF PHASE 2 CLINICAL TRIAL IN AD and PD

### Preliminary data commenced in 1Q2021



A meeting with the FDA to discuss the data from the AD and the PD study as well as from the chronic toxicology in rats and dogs is projected for Fall of 2021



# REVERSAL OF TOXIC CASCADE

## Data from first 14 AD and 14 PD patients

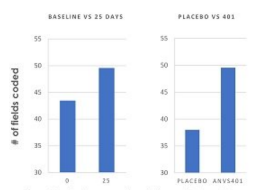
REVERSAL OF TOXIC CASCADE	EXPECTED	ACTUAL OUTCOME	
	OUTCOME	AD	PD
Level of neurotoxic proteins			
Axonal transport	1		
Inflammation			+++
Dead nerve cells	<b>.</b>		
Control proteins	0		
Efficacy: Motor function	1	+	
Efficacy: Cognition	1		+

- +++ p≤ 0.001
- ++ p<u><</u>0.01
- + p≤0.05
- +/- trend
- 0 no change
- opposite result from expected



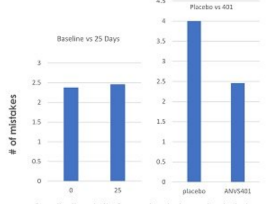
# EFFICACY IN PD PATIENTS - SPEED & COORDINATION

### Data from first 14 PD patients



Speed: Left - the comparison between the treated group with 80 mg/day of ANV\$401 at baseline before treatment and after 25 days on treatment in the rapid coding test. At 25 days the speed is faster than at baseline and they make fewer mistakes (p<0.04).

Right - the comparison between the placebo group and the treated group at 25 days. This graph shows that while the placebo group gets slower, the treated group gets faster (p< 0.04). The lower number shows worse performance.



Coordination: Left - Comparison between treated at baseline and at 25 days. The two scores are identical patients remain stable

Right - the comparison is made between the placebo group and the treated group both at 25 days. The placebo treated group shows a marked deterioration in their motor complications compared to the ANVS401 treated group that was stable (p< 0.07). The lower number shows better performance.



# LOWERING OF INFLAMMATION IN PD PATIENTS

### Data from first 14 PD patients

Inflammatory Marker	% Change from Baseline	p-Value	
Complement C3	-24.9	0.0072	
YKL40	-22.9	0.0213	
sTREM2	-28.2	0.0108	
GFAp	-34.6	0.000001	

The trial measured four inflammatory markers that are prevalent in the brains of AD and of PD patients.

Each of the inflammatory markers showed statistically significant reduction after 25 days of treatment with ANVS401 compared to baseline.



# EFFICACY IN AD PATIENTS - ADAS-Cog11

### Data from 14 AD patients

### Within Data:

From baseline to 25 days in the ANVS401-treated group, ADAS-Cog11 improved by 4.4 points, a statistically significant improvement of 30% (p=0.04).

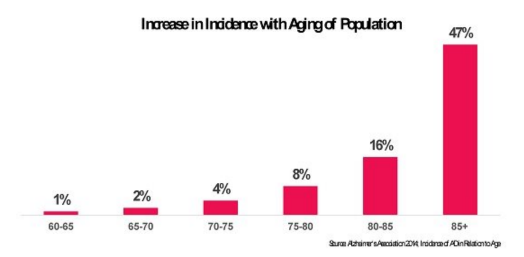
### Between Data:

ANVS401-treated group compared to placebo group at 25 days showed an improvement of 3.3 points, or 22% (p= 0.13).

This is the first double-blind, placebo-controlled study that shows cognitive improvements in AD patients as measured by ADAS-Cog and functional improvements in PD patients as measured by the Unified Parkinson's Disease Rating Scale (UPDRS).



# MARKET PROJECTIONS



Annual sales potential for US and worldwide are over \$100 billion dollars



# Capitalization Table

Common Shares	6,947,269
Options (WAEP: \$2.58)	1,107,598
Warrants (WAEP: \$7.50)	4,800
Fully Diluted Shares Outstanding	8,059,667
Debt	\$0



# MANAGEMENT AND ADVISORY TEAM



### Maria L. Maccecchini, PhD, Founder, President & CEO

Dr. Maccecchini founded Annovis in May 2008 to develop better therapeutics for Alzheimer's, Parkinson's and other neurodegenerative diseases. Was partner and director of two angel groups, Robin Hood Ventures and MidAtlantic Angel Group; Founder and CEO of Symphony Pharmaceuticals/Annovis a biofech company that sold in 2001 to Transgenomic; General Manager of Bachem Bioscience, the US subsidiary of Bachem AG, Switzerland and Head Molecular Biology Mallinckrodt; Dr. Maccecchini did one postdoc at Caltech and one at the Roche Institute of Immunology, her PhD in biochemistry is from the Biocenter of Basel with a two-vear visiting fellowship at The Rockefeller University,



### Jeffrey McGroarty, CPA, MBA, Chief Financial Officer

Mr. McGroarty is a financial executive with experience in investor relations, working with analysts, creditors and financial institutions, planning and analysis, capital allocation, SEC communications and reporting, accounting, acquisitions and tumarounds. He is experienced in effectively managing complex projects, building professional relations and developing staff. Mr. McGroarty was previously employed as CFO of Safeguard Scientifics, Interim Controller at Cephalon, Inc., Vice President-Financial Planning and Analysis of Exide Technologies, Inc., and Senior Manager at PWC, His MBA is from the Wharton School of Business,



### Feng Chang, PhD, VP of Research

Dr. Chang is an experienced neuroscientist with more than a decade of experience in neurodegenerative diseases, with broad scientific knowledge and hands-on experience. Prior to joining Annovis, she was a scientific solution consultant with Clarivate Analytics where she worked on cutting-edge scientific projects with top-50 pharma clients. Previously, Dr. Fang was business development manager for Coriell Institute for Medical Research and an assistant professor at Boston University, where she designed and supervised projects focused on prion diseases and AD as a research feam leader.



### William Mobley, MD, PhD, Chief Scientific Advisor

Distinguished Professor, Department of Neurosciences Florence Riford Chair for Alzheimer Research and Associate Dean for Neurosciences Initiatives at UC San Diego. He is a member of the National Academy of Medicine. His research focuses on the neurobiology of neurotrophic factor actions/signaling and on the hypothesis that malfunction of these mechanisms contribute to neuronal dysfunction in developmental and age-related disorders of the neurosystem.



# SCIENTIFIC ADVISORY BOARD



### Sidney Strickland, PhD, Chairman

Vice President and Dean for Educational Affairs and Research Professor, Patricia and John Rosenwald Laboratory of Neurobiology and Genefics at Rackefeller University, Dr. Strickland's laboratory investigates how dysfunction of the circulatory system contributes to Alzheimer's and other neurodegenerative disorders. He will serve as the Chairman of Annovis Bio's SAB.



### Jeffrey Cummings, MD

Jeffrey Cummings, MD
Dr. Cummings completed Neurology residency and a
Fellowship in Behavioral Neurology at Boston University,
Massachusetts. US training was followed by a Research
Fellowship in Neuropathology and Neuropsychiatry at
he Notional Hospital for Nervous Diseases. London,
England. Dr. Cummings was formerty Professor of
Neurology and Psychiatry, Director of Alzheimer's
Disease Research and Director of the Center for
Neurotherapeutics at UCLA. He was Director of the
Cleveland Clinic Lou Ruvo Center for Brain Health in Las
Vegas Cleveland and Florieta. Vegas, Cleveland and Florida.



### William Mobley, MD, PhD

Dr. Mabley is Distinguished Professor, Department of Neurosciences Florence Riford Chair for Alzheimer Research and Associate Dean for Neurosciences Initiatives at UC San Diego. He is a member of the National Academy of Medicine. His research focuses on the neurobiology of neurotrophic factor actions/signaling and on the hypothesis that malfunction of these mechanisms contribute to neuronal dysfunction in developmental and age-related disorders of the neurosystem.



Dr. Petsko is a member of the National Academy of Sciences, the National Academy of Medicine, the Sciences, the National Academy of Medicine, the American Academy of Arts and Sciences and the American Philosophical Society, His research interests are directed towards understanding interests are directed towards understanding the biochemical bases of neurological diseases like Alzheimer's, Parkinson's, and ALS discovering treatments (especially by using structure-based drug design), that could therapeutically affect those biochemical targets, and seeing any resulting drug candidates tested in humans. He has also made key contributions to the field of protein crystallography.



Rudolph E. Tanzi, PhD
Dr. Tanzi has published over 500 research papers
and has received the highest awards in his field,
including the Metropolitan Life Foundation Award,
Potamkin Prize, Ronald Reagan Award, Silver
Innovator Award, and many others, He was
named to TIME magazine's list of TIME100 Most
Influential People in the World (2015), and
received the Smithsonian American Ingenuity
Award, the top national award for invention and
innovation. He co-authored the popular, trade innovation. He co-authored the popular trade books "Decoding Darkness", New York Times bestseller, "Super Brain", and international bestseller "Super Genes".





# **BOARD OF DIRECTORS**



### Michael B. Hoffman Chairman

Mr. Hoffman is the Founder and Managing Partner of Stone Capital Partners, a private equity firm focused on power and renewable energy. He was Partner of Riverstone, senior managing director at the Blackstone Group and managing director at Smith Barney, Harris Upham & Co. He serves as Chairman of Onconova, Annovis Bio, Curative and is on the Board of Rockefeller University.



### Maria L. Maccecchini, PhD **Executive Board Member**

Dr. Maccecchini founded Annovis in May 2008 to develop better therapeutics for Alzheimer's, Parkinson's and other neurodegenerative diseases. She was the Founder and CEO of Symphony Pharmaceuticals/Annovis, a company focused on protecting brain cells after stroke which was sold in 2001 to Transgenomic.



### Reid S. McCarthy

Mr. McCarthy is experienced in corporate financial management, operations and new venture development. He was CFO of Topaz Pharmaceuticals, Inc. until its sale in 2011 to Sanofi Pasteur. He also served as CFO of JJ Haines & Company, Inc. and provided consulting CFO services to several life sciences companies. He has been a founding executive of several venture capital-backed companies which were successfully sold.

### Claudine E. Bruck, PhD

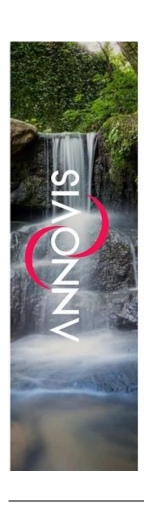
Dr. Bruck is a pharmaceutical executive and scientist with strong entrepreneurial drive. Exhibited successes in building a therapeutic research unit de novo and leading discovery and clinical development of biological (vaccines, biopharmaceuticals) and small molecule medicines as well as an aphthalmic drug portfolio. With creativity and a strong results-focus, she is energized to challenge and lead teams. Extensive Pharmaceutical industry experience spans drug discovery and development across several therapeutic areas.



### Mark White

Mr. White is a biopharmaceutical executive with global marketing, business development and sales experience. Currently, he is an sales experience. Currently, he is an independent consultant and a member of Robin Hood Ventures, a Philadelphia based angel investor group, Frevlously, Mr. White held senior level roles at Pfizer in marketing and commercial level roles at Pitzer in marketing and commercial development, where he led the successful global launches of Inspira, Revatlio, Lyrica and Xeljanz, In his last position, he was Vice President Worldwide Marketing, with global responsibility for new product development and in-line marketing for Pfizer's Inflammation Therapeutic Area.





## INVESTMENT SUMMARY

### A novel approach to treat neurodegeneration is desperately needed

- The markets for AD and PD drugs are in the multibillions of dollars and growing
- Annovis has a novel approach to stop the course of AD and PD
- ANVS401 shows statistically significant improvements in Phase 2a clinical trials:
  - Cognition in AD patients
  - Motor function in PD patients
  - Inflammation in PD patients
- The successful completion of our Phase 2 clinical trials is providing validation of our approach in two diseases and allow us to move to Phase 3 trials in both diseases



# CHANGE IN CAUSES OF DEATH FROM 2000 TO 2018

Breast Cancer - 13%
 Colon Cancer - 21%
 Heart Disease - 21%
 Stroke - 24%
 HIV - 67%
 Parkinson's + 84%

Parkinson's + 84%Alzheimer's + 112%



### ANNOVIS' DRUG ATTACKS MULTIPLE NEUROTOXIC PROTEINS

Chronic and acute brain insults lead to high levels of neurotoxic proteins, to inflammation and neurodegeneration

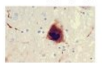
Amyloid B AD/ PD-AB Targeting Compounds



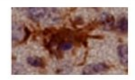
Tau Tauopathies - AD - Tau Targeting Compounds



aSynuclein PD/ AD-aS/NErgeting@mpcurds



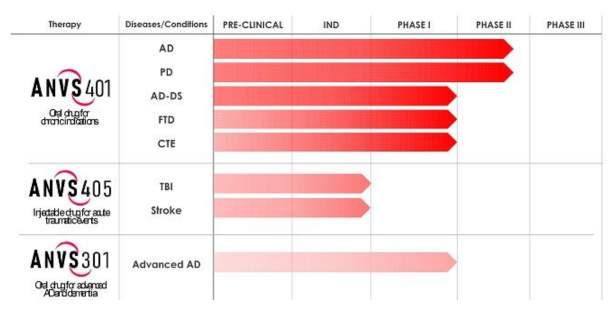
Activated Microglia = High Inflammation



Attacking one neurotoxic protein results in minimal effect ANVS401 is the only drug to attack multiple neurotoxic proteins simultaneously



# **PIPELINE**





# CORPORATE PATENT ESTATE

Multi-layer strategy



Patent/Application	Subject Matter	Status	Expiry
Provisional	ANVS401 to treat viral and bacterial infections of the brain, including Covid19  Pending		2041
PCT	ANVS401 and 405 – Mechanism of Action for prevention and treatment of diseases	Pending	2038
PCT	ANV\$405 - Acute brain and nerve injuries	Granted – Europe and Japan	2036
PCT	ANVS401 - pK/pD, low doses, formulations Neurodegenerative Diseases	Granted – US and Europe	2031
In-licensed patents	Composition of matter, manufacturing, method for treating AD and DS	Granted	2022-25

