Development and Validation of a Wearable Device for Continuous Assessment and Measurement of Sleep-Associated Cranial Fluid **Dynamics and EEG**

Paul Dagum, MD PhD¹, Laurent Giovangrandi, PhD¹, Jacob Winebaum¹, Juan Piantino MD², Miranda M. Lim, MD PhD^{3,4}, Jeffrey Iliff, PhD^{5,6,7} August 17, 2023

¹Applied Cognition, Inc, Redwood City, CA ²Department of Pediatrics, Oregon Health & Science University, Portland, OR ³VA Portland Health Care System, Portland, OR ⁴Department of Neurology, Oregon Health & Science University, Portland, OR ⁵VISN20 Northwest Mental Illness Research, Education and Clinical Center, VA Puget Sound Health Care System, Seattle, WA ⁶Department of Psychiatry and Behavioral Science, ⁷Department of Neurology, University of Washington School of Medicine, Seattle, WA



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DISCLOSURE

- Paul Dagum, MD PhD; Laurent Giovangrandi, PhD; and Jacob Winebaum ulletare employed by Applied Cognition and own stock in the company
- Jeffrey Iliff, PhD and Miranda Lim, MD PhD receive compensation from ulletApplied Cognition and hold stock options in the company

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Glymphatic Flow: A New Biology of the Brain The newly discovered glymphatic system plays a critical role in the clearance of neurodegenerative

The newly discovered glymphatic system plays a critical role in the clearant proteins and metabolic waste products



Slow Wave Activity

The cleaning power of **slow wave activity (SWA**) during deep sleep is augmented by a 60% increase in the interstitial fluid (ISF) volume created via AQP-channels.

Arterial Pulsatility

Arterial pulsatility in the brain provides the motive force that moves cerebrospinal fluid (CSF) into the perivascular spaces surrounding major arteries.





Pulsatile Waves of CSF

Pulsatile waves of CSF flow increase waste clearance and are entrained to restorative slow wave oscillations by hemodynamic oscillations and **neurovascular coupling**. **Glymphatic System:** a waste clearance pathway in the brain that relies on interchange of cerebrospinal fluid (CSF) and interstitial fluid (ISF).

Ne

Para-arterial space

AQP4 ____

Astrocyte vascular endfeet





What we know about glymphatic flow: from pre-clinical to clinical Pre-clinical studies have demonstrated that this new biology has profound effects on brain health and

disease

SCIENCE ADVANCES | RESEARCH ARTICLE

NEUROPHYSIOLOGY

Increased glymphatic influx is correlated with high EEG delta power and low heart rate in mice under anesthesia

Lauren M. Hablitz¹, Hanna S. Vinitsky¹, Qian Sun¹, Frederik Filip Stæger², Björn Sigurdsson², Kristian N. Mortensen², Tuomas O. Lilius^{2,3}, Maiken Nedergaard^{1,2}*



- Single measure per rodent before sacrifice
- Different anesthetic protocols
- EEG measured in a different population of rodents



Repetitive Mild Traumatic Brain Injury Alters Glymphatic Clearance Rates in Limbic Structures of Adolescent Female Rats

Jennaya Christensen^{1,2}, David K. Wright², Glenn R. Yamakawa², Sandy R. Shultz^{2,3} & Richelle Mychasiuk^{1,2,4,5*}

Brain-wide glymphatic enhancement and clearance in humans assessed with MRI

Geir Ringstad,^{1,2} Lars M. Valnes,³ Anders M. Dale,^{4,5,6} Are H. Pripp,⁷ Svein-Are S. Vatnehol,⁶ Kyrre E. Emblem,⁹ Kent-Andre Mardal,^{3,10} and Per K. Eide^{2,11}



Figure 3. Scatter plot with bar graphs displaying the one-way ANOVA results for the time required to reach maximum signal intensity, a measure of glymphatic influx, for the (A) amygdala, (B) hippocampus, (C) hypothalamus, and (D) olfactory bulb. Time to max is calculated from the start of contrast agent injection. Means \pm standard error; * main effect of injury, p < 0.05.

- Animal model of rmTBL ٠
- Showed impact on glymphatic influx/efflux ٠



duration of the study

Intrathecal injection of Gadobutrol Serial MRIs over 48 hours with participant supine for the

Measurement of glymphatic function using iv contrast-enhanced MRI

The current gold standard measurement of glymphatic flow in humans requires an iv bolus of contrast followed by serial MRIs



The lack of continuous in-human measurement of glymphatic function is limiting our understanding of this transformative biology and its potential in therapeutic discovery

Richmond et al. Eur J Neurosci 2023

The Science of How We Measure Glymphatic Flow*

С

Our direct measurement of glymphatic flow measures the change in brain parenchymal resistance (or flow at a given pressure gradient) using continuous spatial measurements of conduction through the brain at different frequencies







- Slow wave activity
- Arterial pulsatility
- Pulsatile waves of CSF

Network models from Nedergaard's group reveal that change in brain resistance in sleep is the key driver to glymphatic flow

Article

A network model of glymphatic flow under different experimentally-motivated parametric scenarios

Jeffrey Tithof,^{1,2,5,*} Kimberly A.S. Boster,¹ Peter A.R. Bork,³ Maiken Nedergaard,^{3,4} John H. Thomas,¹ and Douglas H. Kelley







*Dagum et al Patent US20230080140A1

Our device sensors also measure key physiologic drivers of glymphatic flow using novel approaches to radically miniaturize and simplify instrumentation, allowing for continuous overnight measurements



Glymphatic System:

a waste clearance pathway in the brain that relies on the interchange of cerebrospinal fluid (CSF) and interstitial fluid (ISF)

Benchmarking Study: Primary Objective

To demonstrate that our investigational device is as good or better at **measuring glymphatic function** than gold-standard neuroimaging and can do so continuously during sleep which is currently not possible



Clinical studies **completed in June**:

- Benchmarking study is a 30 participant randomized cross-over design at the University of Florida
 - Compares our device to CE MRI neuroimaging of glymphatic function
 - Identifies neurophysiology and blood biomarker correlates with device measurements
- Replication study is a comparable 16 participant randomized cross-over design at the University of Washington



Cognitive Testing Contrast Injection

Study Results: First-ever Continuous Measure of Glymphatic **Flow in Humans**

Participants who slept had lower brain resistance to glymphatic flow (p < 0.0005)



Significant Study Findings

- movement into the brain with sleep
- 0.018)

Resistance (p= 0.01, 0.011, 0.039, 0.044), not sleep/wake assignment (three p=NS, one p = 0.025), was the significant moderator of MRI contrast

> Changes in resistance correlated with changes in sleep EEG band power replicating pre-clinical animal studies (all p < 0.005 except beta p =

Changes in overnight resistance (p = 0.005), not sleep/wake assignment (p=NS), predicted performance in morning cognitive test battery

Change in resistance was a robust predictor of contrast movement from the blood and CSF compartments through the brain parenchyma*

Moderation of Solute Transport from Blood to Brain by

Overnight Flow Resistance

Lower resistance steepened the relationship between CSF-parenchymal contrast (glymphatic influx) and flattened the relationship between blood-parenchymal contrast (clearance to blood)

	Moderation of Morn	ing Brain Solute Transport by Ove	rnight Resistance
Predictors	Estimates	CI	р
(Intercept)	0.4074	-0.0167 - 0.8315	0.063
Overnight Resistance	-1.2833	-2.51950.0471	0.044
Morning CSS Contrast	0.3372	-3.5942 - 4.2685	0.868
Morning Ventricular Contrast	-0.5336	-0.97210.0952	0.018
Resistance:CSS	8.3833	1.9704 – 14.7962	0.011
Resistance: Ventricles	1.3555	0.0839 - 2.6272	0.039
CSS:Ventricles	1.9429	-1.9562 - 5.8419	0.333
Resistance:CSS:Ventricles	-8.8356	-15.44292.2283	0.010
Random Effects			
σ^2	0.001		
$\tau_{00 pid}$	0.002		
ICC	0.632		
N _{pid}	29		
Observations	357		





Moderation of Morning Solute Transport by Overnight Flow Resistance

9

Brain parenchymal resistance is dynamically coupled to sleep neurophysiology in humans

We used *continuous device recordings* of resistance, EEG and cardiovascular metrics during sleep to identify coupling between sleep neurophysiology and resistance



Data was time aligned in the device to within 4

Device sleep hypnogram was used to identify

EEG power bands (delta, theta, alpha and beta), HR and HRV (sdnn) were computed

First difference of all measures were taken (R,

Linear, multilinear and linear threshold models

Delta Power Decreased Resistance in NREM Sleep

Delta power showed a threshold effect on resistance requiring a step increase > 7%. A step increase in delta power had a larger effect on R than a similar size step decrease

	Change in Brain Parenchymal Resistance with Threshold Increase Above 7% in Delta Power During NREM Sleep			
Predictors	Estimates	CI	р	
(Intercept)	506.4601	157.0480 - 855.8723	0.005	
Delta Power %	-52.7240	-87.565017.8830	0.003	
Observations	90			
R^2/R^2 adjusted	0.093 / 0.083			

	Change in Brain Parenchymal Resistance with Decrease Below Zero in Delta Power During NREM Sleep			
Predictors	Estimates	CI	р	
(Intercept)	-110.9821	-166.717455.2468	<0.001	
Delta Power %	-17.6942	-28.34887.0397	0.001	
Observations	708			
R^2/R^2 adjusted	0.015 / 0.013			

Does this explain why morning brain parenchymal resistance does not return to evening levels?



Change in Brain Parenchymal Resistance with Threshold Increase in Delta Power During NREM Sleep

tial. Do not distribute. 11

Beta Power Increased Resistance in NREM Sleep

A step increase/decrease in beta power led to a step increase/decrease in resistance during NREM sleep

Change in Brain Parenchymal Resistancewith with Change in Beta Power During NREM Sleep CI **Predictors** Estimates р 0.025 -29.8051 -55.8753 - -3.7348(Intercept) Beta Power % 14.8749 2.5577 - 27.19220.018 Observations 1402 0.004 / 0.003 R^2 / R^2 adjusted

Change in Brain Parenchymal Resistance with Change in Beta Power During NREM Sleep





HR and HRV had Opposite Effects on Resistance During NREM Sleep

Both where threshold above zero with a step increase in HR leading to a step increase in R and a step increase in HRV leading to a step decrease in R

	Change in Brain Parenchymal Resistance with Threshold Increase Above 0.28bpm in HR During NREM Sleep			
Predictors	Estimates	CI	р	
(Intercept)	-98.1940	-165.837830.5502	0.005	
HR	27.5603	0.2205 - 54.9001	0.048	
Observations	450			
R^2 / R^2 adjusted	0.009 / 0.006			

- 004 - 005 - 005

Change in Brain Parenchymal Resistance with Threshold Increase Above 34 ms in HRV During NREM Sleep

Predictors	Estimates	CI	р
(Intercept)	462.0403	170.2266 - 753.8540	0.002
HRV	-8.4560	-13.07703.8349	<0.001
Observations	86		
_ 2 2	0 126 / 0 126		

 R^2 / R^2 adjusted 0.136 / 0.126

Change in Brain Parenchymal Resistance with Increase in HRV During NREM Sleep

5.0 Change in HR

2.5



NREM Sleep





Theta Power Decreased and Beta Power Increased Resistance in REM Sleep

A step increase (decrease) in theta power (beta power) during REM sleep led to a step decrease (increase) in resistance





Why Do We Care? Lower overnight resistance maintained sleep-sensitive cognitive performance

Participants who had lower overnight resistance, or greater glymphatic flow, performed better on a multi-domain cognitive battery in the morning

	Predictors of Cognitive Standardized Test Scores			
Predictors	Estimates	CI	р	
(Intercept)	4.36	2.40 - 6.31	<0.001	
Overnight Mean Resistance	-1.81	-2.710.91	<0.001	
HRV LF/HF Ratio	-0.14	-0.220.07	<0.001	
Age [yrs]	-0.04	-0.070.01	0.010	
Random Effects				
σ^2	0.31			
τ _{00 Cog.Test}	0.22			
$\tau_{00 pid}$	0.61			
N Cog.Test	5			
N _{pid}	44			
Observations	340			
Marginal R^2 / Conditional R^2	0.173 / NA			







Lower overnight resistance was also weakly associated with greater β amyloid clearance from the brain

Overnight differences in serum Aβ40 and Aβ42 tended to be greater with lower overnight resistance

	Predictors of Overnight Change in Blood Abeta42		
Predictors	Estimates	CI	р
(Intercept)	7.81	-4.54 - 20.15	0.211
Overnight Mean Resistance	-9.48	-21.80 - 2.84	0.129
Sleep Visit	-1.96	-3.870.04	0.045
Observations	70		
R^2 / R^2 adjusted	0.066 / 0.038		

	Predictors of Overnight Change in Blood Abeta		
Predictors	Estimates	CI	p
(Intercept)	97.47	-8.77 - 203.71	0.072
Overnight Mean Resistance	-103.99	-209.96 - 1.98	0.054
Sleep Visit	-13.97	-30.54 - 2.60	0.097
Observations	71		
R^2/R^2 adjusted	0.066 / 0.039		



Research Roadmap

With a non-invasive investigational device to measure glymphatic function in the field, we will better understand the pathophysiology and clinical correlates of glymphatic dysfunction to restore cognitive performance and improve injury recovery

Pathophysiology

- Identify the impact of sustained or repeated exposure to extreme environments and trauma
- Identify the impact of diseases such at TBI and mental health on glymphatic function

Clinical Performance Correlates

Identify the sleep, cognitive and neurological performance correlates of glymphatic pathophysiology

Therapeutic Discovery

Investigate pharmacological and non-pharmacological interventions to restore glymphatic function