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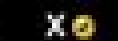
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Kristi Hines

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@Twitter · Jul 21

We've had tremendous interest in the new ads revenue sharing program from creators around the world.

Creators who are eligible will automatically be able to start receiving ads revenue upon launch — all you need to do is set up payouts and you'll be good to go!

Ads Revenue

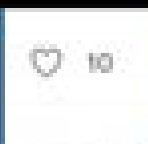
@Twitter · Jul 13

Surprise! Today we launched our Creator Ads Revenue Sharing program.

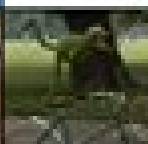
We're expanding our creator monetization offering to include ads revenue sharing for creators. This means that creators can get a share in ad revenue, starting in the replies to their posts. This is part of



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Messages

X App This Month

SJ Ball

X App This Month:

Monthly Weather Review ,1987 *The Propagation of Neurodegenerative Diseases by Inflammation and Exosomes*

Valerie Sackmann, 2019-10-16 Alzheimer's disease AD and Parkinson's disease PD are the two most common neurodegenerative diseases with rates increasing along with the ageing global population. Despite best efforts we still do not understand the etiopathogenesis of these diseases and there are no effective disease modifying treatments. Cognitive deficiencies or motor complications that emerge during AD and PD are thought to be the result of the accumulation of misfolded aggregate prone proteins such as amyloid A and tau or synuclein syn respectively. Growing evidence suggests that prefibrillar oligomers of A and syn oA and o syn are key contributors to the progression of these diseases. The progressive accumulation of these proteins leads to a gradual spread of pathology throughout interconnected brain regions but the mechanisms by which this spreading occurs are still largely unknown. Neuroinflammation has been recognised as an important contributor to neurodegenerative disease. It is hypothesised that a pro inflammatory environment initiated by the innate immune system either through activation from A itself or indirectly through neuronal injury signals in AD. These phenomena are thought to either cause or accelerate AD such that an anti inflammatory approach may be neuroprotective. In paper I we investigated whether different inflammatory environments affected the transfer of oA between neuron like cells in addition to investigating inter and intracellular protein changes. This study demonstrated that an anti inflammatory environment reduces the transfer of oA between cells. We also provide evidence that these cells begin to take on the phenotype of the inflammatory milieu while also demonstrating that the expression profile of endosomal lysosomal and protein trafficking proteins is altered during these conditions. Small extracellular vesicles called exosomes which are key players in cell to cell communication have been proposed to play an influential role in spreading neurodegenerative proteins between cells. Exosomes are small membranous vesicles that are formed by the inward budding of multivesicular bodies (MVBs). These MVBs can then merge with the plasma membrane to be released into the extracellular environment as vesicles which serve as vehicles for transferring proteins, lipids and mRNAs between cells. The ESCRT dependent pathway is the most understood mechanism underlying exosome biogenesis. However, exosomes can also be formed through ESCRT independent pathways including through the hydrolysis of sphingomyelin by neutral sphingomyelinase 2 (nSMase2) which produces ceramide. Paper II investigated whether exosomes formed through an ESCRT independent pathway plays a significant role in the transfer of o syn between neuron like cells. As oxidative stress is a common feature in PD brains which in turn dysregulates nSMase2 activity we also tested our model under hypoxic conditions. Inhibition of nSMase2 significantly reduced the transfer of o syn between cells but also resulted in decreased syn aggregation. Hypoxia did not influence o syn transfer however it significantly dysregulated the sphingolipid composition which may be important for syn binding to exosomes and exosome communication. During AD and PD there is a noted reduction in the effectiveness of autophagy a

process critical to cellular proteostasis Recent studies have uncovered shared regulatory mechanisms of exosome biogenesis and autophagy suggesting that they are closely linked Previous findings have shown that inhibition of autophagy in AD mice mediates A trafficking through altering the secretion of A in MVBs To further study this effect we investigated the interplay between autophagy and exosome secretion using ATG7 knock out x APPNL F knock in AD mice in paper III These autophagy deficient AD mice had a reduced extracellular A plaque load but increased intracellular A which was found to be assembled into higher ordered assemblies While exosomal secretion was dysregulated in these mice the amount of A packaged into the exosomes was unchanged Lastly one of the biggest challenges in developing effective treatments for AD is the lack of early diagnosis of living patients As the connection between exosomes and the spread of neurodegenerative proteins is still relatively new there remains a diagnostic potential to be explored with exosomes Paper IV aimed to develop a new diagnostic assay to detect oA in exosomes isolated from human cerebrospinal fluid Although exosomal oA was readily detected in some of these samples the assay's sensitivity requires additional optimisation before it can be further validated for the clinic In summary the studies presented in this thesis have furthered our understanding of how inflammation autophagy and exosomes contribute to the intercellular transmission of AD and PD associated proteins We have shown that an anti inflammatory approach may slow down the progression of AD through reducing the transfer of oA between cells We also provide novel findings relating to the biogenesis of exosomes which in turn affected the ability of exosomes to transmit neurodegenerative proteins between cells and their association with autophagic processes Finally we have investigated the feasibility of exosomes as an early AD diagnostic marker This work has helped to elucidate some of the mechanisms underlying the progression of neurodegenerative diseases which may be useful targets for the investigation of new therapeutic avenues

The National Teachers' Monthly, 1877 **The American Mathematical Monthly**, 1900 Includes section Recent publications **Sidereal Messenger, a Monthly Review of Astronomy**, 1889 **Graphic Arts Monthly and the Printing Industry**, 1954 *The later legislation of the Pentateuch* John William Colenso, 1871 **Monthly Foreign**

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Advertising and Press Annual of Africa, 1952 *Docket No. 73-1957*, 1974 **Barclays Law Monthly**, 1983-07
Records & Briefs New York State Appellate Division, *Cassier's Engineering Monthly*, 1911 **California. Court of Appeal (1st Appellate District). Records and Briefs** California (State)., Number of Exhibits 3 **Monthly Notices of the Royal Astronomical Society**, 1885

Decoding **X App This Month**: Revealing the Captivating Potential of Verbal Expression

In a time characterized by interconnectedness and an insatiable thirst for knowledge, the captivating potential of verbal expression has emerged as a formidable force. Its ability to evoke sentiments, stimulate introspection, and incite profound transformations is genuinely awe-inspiring. Within the pages of "**X App This Month**," a mesmerizing literary creation penned with a celebrated wordsmith, readers attempt an enlightening odyssey, unraveling the intricate significance of language and its enduring impact on our lives. In this appraisal, we shall explore the book's central themes, evaluate its distinctive writing style, and gauge its pervasive influence on the hearts and minds of its readership.

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