

higher in the ICI cardiotoxicity group ( $6.8 \pm 4.2$  vs.  $1.0 \pm 1.7$ ,  $p=0.017$ ). The proportion of abnormal myocardium was higher in the ICI cardiotoxicity group ( $66 \pm 21\%$  vs.  $45 \pm 18\%$ ,  $p=0.050$ ), as well as the proportion of myocardium found to be dysfunctional ( $26 \pm 22\%$  vs.  $3.0 \pm 6.0\%$ ,  $p=0.041$ ). **DISCUSSION/SIGNIFICANCE OF FINDINGS:** Despite having preserved LVEF, patients who met criteria for ICI-associated cardiotoxicity had both global and regional abnormal LV strain. Fast-SENC imaging may provide a sensitive tool for detection of early cardiotoxicity in this population. This study is limited by its small cohort and a larger prospective study would be of value.

65993

### Peptide Conjugated Hollow, Degradable Nanoparticles Bind to Exposed Hyaluronic Acid for the Prevention and Treatment of Osteoarthritis

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**ABSTRACT IMPACT:** Our research would be the first therapeutic to both prevent and treat osteoarthritis - helping 27 millions U.S. citizens alone immediately. **OBJECTIVES/GOALS:** Our objective is to conjugate hyaluronic acid binding peptides (HABP) to anionic hollow nanoparticle (hNP), and allowing the HABP-hNP complex to penetrate into osteoarthritic cartilage, bind to exposed HA, prevent further degradation, and restore the compressive strength of articular cartilage. **METHODS/STUDY POPULATION:** N-isopropyl acrylamide, 2-acrylamido-2-methyl-1-propanesulfonic acid, N,N'-bis(acryloyl)cystamine, and Acrylic Acid, in fluorescent batches rhodamine b isothiocyanate (RBITC), were polymerized via precipitation reaction. HA binding peptide, GAHWQFNALTVRGSG-Hydrazide (GAH-Hyd), was covalently bonded to the hNP using DMTMM chemistry. The reaction was halted by diluting the solution 10:1 with milliQ water and purified using tangential flow filtration. The dynamic viscosity of the six treatments were analyzed in a 70 kDa HA. Using a rheometer (Discovery HR-3) with a 20 mm parallel plate geometry, TA Instruments, New Castle, DE), a frequency sweep (0.01 -1000 Hz, 2.512 Pa) was conducted to measure the storage modulus of each solution. **RESULTS/ANTICIPATED RESULTS:** GAH-Hyd was successfully conjugated to the surface of the hNP and zeta-potential shows a significant increase in surface charge from -21.41 mV for unconjugated hNP to -8.94 mV for 65 GAH conjugated hNP, confirming conjugation. The hNPs need  $65 \pm 10$  GAH per nanoparticle to significantly bind to HA, shown by increasing the dynamic viscosity of the solution. The minimum concentration of 65 GAH-hNP required to significantly bind to HA is  $313 \mu\text{M}$ . These data from our study display the ability to functionalized the surface of polymeric hNPs with site specific peptides and their ability to bind to diseased tissue. We expect the GAH-hNP system will restore the compressive strength of OA cartilage and prevent further HA degradation in ex vivo aggrecan depleted cartilage plugs. **DISCUSSION/SIGNIFICANCE OF FINDINGS:** Binding to exposed HA within the ECM of cartilage protects the HA from further degradation, halting the progression of OA. 65 GAH-hNP binds to HA at a  $313 \mu\text{M}$ . Our system can be translated and used to treat a multitude of conditions by conjugating tissue specific peptides to the surface of our hNPs and delivery site specific therapeutics to diseases tissue.

## Regulatory Science

19751

### Identifying Barriers to Diabetes Technology in Low-Income, Type 1 Patients

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**ABSTRACT IMPACT:** This research will aid clinical and policy solutions on lessening the vast health disparities and overall access issues for low-income, type 1 diabetes patients. **OBJECTIVES/GOALS:** Identify key barriers to accessing continuous glucose monitors (CGMS) and care options for low-socioeconomic status (SES) patients on public insurance. Low-SES patients with type 1 diabetes (T1D) have lower utilization rates of effective diabetes management technologies and worse clinical outcomes. **METHODS/STUDY POPULATION:** A literature review was conducted to understand the current research landscape for T1D and lead to the identification of potential barriers which included socioeconomic status, low-income, health literacy, and racial/ethnic minority. Clinicaltrials.gov was searched using the keyword 'type 1 diabetes' in conjunction with the identified barriers (as well as the keyword 'barrier'). A follow up review of each state's Medicaid programs was conducted to analyze cost and access options for CGMs and the overall financial burden of the disease on low-SES T1D patients. States that offered CGM coverage were further analyzed to determine reimbursement rates and actual out-of-pocket cost for patients. **RESULTS/ANTICIPATED RESULTS:** Of 285 trials identified from Clinicaltrial.gov searches, only seven relevant trials examined barriers and T1D for low-SES patients. Additionally, many of these studies, both in and outside of the clinical trial space, seldom distinguished between type 1 and type 2 diabetes" an important distinction given that T1D has a higher financial burden and a quicker onset of complications. Currently, 39 states offer various insurance coverage through their Medicaid programs, but have clinical restrictions and requirements such as pediatric coverage only or minimum blood glucose requirement checks. Additionally, there is vast variability in reimbursement rates between states (\$0-\$800). **DISCUSSION/SIGNIFICANCE OF FINDINGS:** Study results indicate less effective diabetes management for low-SES T1D patients and a need for more intersectional clinical trial research. Differences in state's Medicaid CGM coverage, expressed in disparate clinical outcomes for these T1D patients, belies financial incentives to health improvements, as annual US T1D costs are \$14.4 billion.

67702

### At-Home Screening Tool for Anosmia

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**ABSTRACT IMPACT:** By developing and validating a simple and cost-effective at-home screening tool for loss of smell, we can efficiently detect infection with COVID-19, neuropsychiatric disease such as Alzheimer's, and post-operative smell loss. **OBJECTIVES/GOALS:** To develop and validate a feasible and cost-effective screening tool for olfactory dysfunction (OD) using common household items. **METHODS/STUDY POPULATION:** The study has two phases. In the Development phase, 120 participants with