Drug Loading Capacity of Environmentally Sensitive Polymeric Microgels

RYAN MCDONOUGH, KIRIL STRELETZKY, MEKKI BAYACHOU, PUBUDU PEIRIS, Cleveland State University — Microgel nanoparticles consisting of cross-linked polymer hydroxypropyl cellulose chains have a temperature dependent volume phase transition, prompting the use of microgels for controlled drug transport. Drug particles aggregate in the slightly hydrophobic interior of microgels. Microgels are stored in equilibrium until the critical temperature ($T_v$) is reached and the volume phase transition limits available space, thus expelling the drugs. Our study was designed to test this property of microgels using amperometric electrochemical methods. A critical assumption was that small molecules inside microgels would not interact via diffusion with the electrode surface and thus total current would be decreased across the electrodes in a microgel sample. A room temperature ($T_{room}$) flow amperometric measurement comparing microgel/tylenol solution with control tylenol samples yielded about 20% tylenol concentration reduction of the microgel sample. Results from the steady state electrochemical experiment confirm the presence of about 20% tylenol concentration drop of the microgel sample compared to control sample at $T_{room}$. Using the steady-state experiment with a cyclic temperature ramp from $T_{room}$ to beyond $T_v$ showed that the tylenol concentration change between the temperature extremes was greater for the microgel solution than for the control solution.

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