Steffen Vowinkel<sup>1</sup>, Stephen Paul<sup>2</sup>, Torsten Gutmann<sup>2</sup> and Markus Gallei<sup>1,\*</sup>

Supporting Information



Figure S1. TEM Images of P(MMA-co-ALMA)@P(EA-co-MPSIsoprop) particles.



Figure S2. TEM Images of P(MMA-co-ALMA)@P(EA-co-MPSMeEt) particles.





Nanomaterials 2017, 7, 155; doi: xxxxxxxxx

www.mdpi.com/journal/nanomaterials



Figure S3. TEM Images of P(S-co-ALMA)@P(EA-co-MPSEt) particles - 230 nm (DLS).

Figure S4. TEM Images of P(S-co-ALMA)@P(EA-co-MPSEt) particles - 289 nm (DLS).



**Figure S5.** DLS measurement of different sized P(S-*co*-ALMA) particles with corresponding shell of P(EA-*co*-MPSEt).





**Figure S7.** Possibilities of cross-linking of the 3 methacryloxypropyltriethoxysilane (MPSEt) linker indicated by  $T_n$  groups (n=0, 1, 2, 3) in the <sup>29</sup>Si CP MAS spectra. Note that cross-linking between different linker molecules L is more probable than the binding to bulk silica. The obtained bulk silica indicated by  $Q_n$  groups (n= 1, 2, 3) in the <sup>29</sup>Si CP MAS spectra is assumed to be formed by decomposition of the organic silane linker.



**Figure S8.** <sup>13</sup>C static spectrum of the free linker MPSEt with signal assignments, and corresponding <sup>13</sup>C CP MAS spectrum measured at 10 kHz spinning of the lyophilized powder sample prepared with this linker.



**Figure S9.** <sup>13</sup>C static spectrum of the free linker MPSIsoprop with signal assignments, and corresponding <sup>13</sup>C CP MAS spectrum measured at 10 kHz spinning of the lyophilized powder sample prepared with this linker.



**Figure S10.** <sup>13</sup>C static spectrum of the free linker MPSMeEt with signal assignments, and corresponding <sup>13</sup>C CP MAS spectrum measured at 10 kHz spinning of the lyophilized powder sample prepared with this linker.



**Figure S11.** <sup>29</sup>Si CP MAS spectra of samples based on the MPSIsoprop linker system in three different states: (a) lyophilized powder, (b) hybrid film and (c) hybrid film after heat treatment, and signal assignment of  $T_n$  groups (n=0,1,2,3) and  $Q_n$  groups (n=1,2,3). Spectra were measured at 14 T at a spinning rate of 8 kHz.



**Figure S12.** <sup>29</sup>Si CP MAS spectra of samples based on the MPSMeEt linker system in three different states: (a) lyophilized powder, (b) hybrid film and (c) hybrid film after heat treatment, and signal assignment of  $T_n$  groups (n=0,1,2,3) and  $Q_n$  groups (n=1,2,3). Spectra were measured at 14 T at a spinning rate of 8 kHz.



**Figure S13.** SEM images of the surface (left) and cross-section (right) of the P(MMA-*co*-ALMA)@P(EA*co*-MPSEt) film after processing at 140°C and 180 bar for 3 minutes.



**Figure S14.** SEM images of the surface (left) and cross-section (right) of the P(MMA-*co*-ALMA)@P(EA*co*-MPSIsoprop) film after processing at 140 °C and 180 bar for 3 minutes.



**Figure 15.** SEM images of the surface (left) and cross-section (right) of the P(MMA-*co*-ALMA)@P(EA*co*-MPSMeEt) film after processing at 140 °C and 180 bar for 3 minutes.