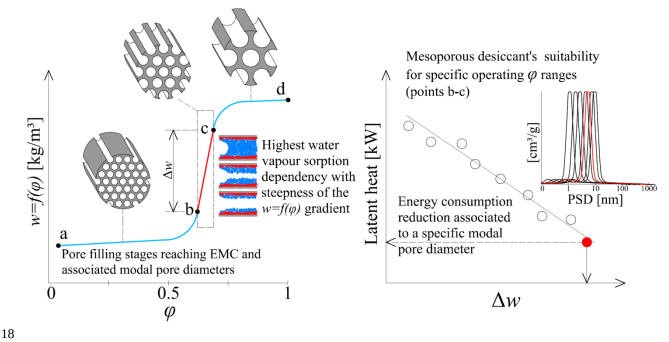
1 Hygrothermal simulation-informed design of mesoporous desiccants for optimised

2 energy efficiency of mixed mode air conditioning systems

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#### 21 Abstract

This paper describes an optimization technique using hygrothermal numerical modelling to 22 determine an ideal and unknown isotherm in order to inform the design of optimised 23 24 mesoporous desiccants. Their suitability for passive humidity buffering as well as their impact on energy efficiency was assessed when assisting a mixed mode air-conditioning (AC) 25 system. Three clear stages of water vapour adsorption were found that strongly correspond to 26 27 the  $\Delta w$  gradient when assessing the kinetics of adsorption and exchange rates for periodic moisture loads. Consistent agreement was found between the latent heat of dehumidification 28 used by the AC system and the desiccant decay time after successive sorption loop cycles. 29 This confirmed the material's suitability for specific applications and was found to be highly 30 sensitive to the portion of the isotherm between  $\varphi_{i,L}$  -  $\varphi_{i,U}$  ( $\Delta w$  gradient), compared with full 31 adsorption capacity (total w) when assessing total energy consumption. The experimental 32 results of sorption kinetics appeared to be slightly underestimated between the  $\Delta w$  gradient 33 and the response time to reach equilibrium moisture content (EMC). The major 34 35 underestimations were found to be consistent with the kinetics of adsorption/desorption when analysing their significance based on w differences. These were largely attributed to a 36 combination of adsorption kinetics (time-response) and adsorption/ desorption hysteresis. 37 However, this was not evident when comparing long-term experimental data and numerical 38 estimations for water vapour sorption isotherms, since numerical model accurately predicted 39 them. This suggests that both adsorption kinetics and the scanning curve prediction, within a 40 hysteresis loop, are not accurately represented by current hygrothermal models and are hence 41 a priority for future research. 42

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Keywords: hygrothermal modelling; water vapour isotherm; desiccants design; mesoporous;
air conditioning; energy efficiency

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# 49 Nomenclature

50	ρ	Bulk density	kg/m³
51	Cp	Specific heat capacity	J/(kg K)
52	$\lambda(w)$	Moisture-dependent thermal conductivity	W/(m K)
53	φ	Relative vapour pressure $(p_v/p_{sat})$	-
54	n	Bulk porosity	m³/m³
55	$w = f(\varphi)$	Water vapour sorption isotherm	kg/m³
56	$\mu(\varphi)$	Water vapour diffusion factor	-
57	$D_w(w)$	Liquid water diffusivity	m²/s
58	RH	Relative humidity; where $RH = \varphi \cdot 100$	%
59	$p_{\nu}/p_0$	Partial pressure of water vapour	-
60	Н	Total enthalpy	J/m <sup>3</sup>
61	$h_{v}$	Latent heat of phase change	J/kg
62	t	Time	S
63	Т	Temperature	Κ
64	$\delta_p$	Vapour permeability	kg/(m s Pa)
65	Suffixes:		
66	i	interior	
67	е	exterior	
68	L	lower limit	
69	U	upper limit	
70			
71			

### 72 **1 Introduction**

Desiccant material performance is typically assessed with respect to isotherm characteristics 73 and target application <sup>1, 2</sup>. This implies that the criteria for optimisation are dependent upon 74 75 the thermal and moisture loads, and operating parameters, inside the closed environment being studied. Optimised mesoporous desiccants can be used to passively regulate indoor dry 76 bulb temperature and relative humidity in closed environments, and to reduce the energy 77 penalty of mechanical heating, ventilation and air conditioning (HVAC)<sup>3,4</sup>. Traditional 78 refrigerant vapour compressor systems (VCS) used for regulating the interior psychrometric 79 conditions of 'closed environments' (e.g., offices, hospitals, laboratories, industry, etc.) are 80 responsible for a large amount of energy consumption with associated greenhouse gas 81 emissions 5-8. The study of desiccants as dehumidifier materials in the air-conditioning (AC) 82 industry has attracted the interest of many researchers in the last decades <sup>9-13</sup>. This has 83 primarily motivated the understanding their abilities to reduce the 'energy penalties' caused 84 by mechanical air dehumidification (latent heat), and the required energy to reheat the air up 85 to the supply air temperature (sensible heat), by engineering their water adsorption/retention 86 properties <sup>14-17</sup>. Some examples of desiccant implementation are included in (i) AC mixed 87 mode systems <sup>18, 19</sup>, (ii) evaporative cooling technology for high humidity climates <sup>20, 21</sup>, (iii) 88 regulating indoor thermal comfort <sup>22-24</sup>, (iv) as a solid–desiccant in solar AC systems <sup>25-28</sup>, (v) 89 indirect evaporative cooling systems <sup>29, 30</sup>, and (vi) waste heat transformation applications for 90 absorption chillers and heat pumps <sup>31-34</sup>. 91

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As an alternative to traditional VCS cooling technologies, solid–desiccant cooling systems
(SDC) have motivated the investigation of advanced desiccant materials since their
characteristics (water adsorption behaviour, regeneration capacity, and long–term stability)
play an important role in their performance and efficiency <sup>35, 36</sup>. These have been organized
into three main categories <sup>37</sup>: (i) composite desiccants (*i.e.*, silica-based, mesoporous silicate-

based and carbon-based), (ii) nanoporous inorganic materials (*i.e.*, aluminosilicate zeolites, 98 aluminophosphates, and aerogels), and (iii) polymeric desiccants (*i.e.*, metal–organic 99 frameworks (MOFs) and polyelectrolyte). Ongoing research <sup>38-42</sup> has been focused on the 100 101 preparation of these materials with improved water adsorption characteristics which include high surface area, large pore volume and an unprecedented physicochemical variability. This, 102 in the case of MCM-41 and SBA-15, has been possible due to the modulation of their textural 103 properties through the synthesis process by controlling temperature <sup>43, 44</sup> and reaction time <sup>45,</sup> 104 <sup>46</sup>. These mesoporous materials exhibit enhanced water adsorption properties compared with 105 those traditional silica-based as a result of having regular pore structures and high surfaces 106 107 areas <sup>47</sup>. On the other hand, significant interest has been focussed on MOFs as water adsorbents due to their large adsorption capacity, fine-tunable pore size, high surface area, 108 and pore volume controlled by change in organic ligands and metal sub–units <sup>48-51</sup>. In recent 109 work done by Furukawa *et al.*<sup>52</sup>, large water adsorption capacity and pore volume (735–850) 110 cm<sup>3</sup> g<sup>-1</sup> and 0.88 cm<sup>3</sup> g<sup>-1</sup>, respectively) was reported from low to high  $p_{y}/p_{0}$  range (0.05–0.7). 111 Their MOF materials exhibited outstanding sorption/desorption water uptake after 5 cycles 112 (showing no loss of porosity) at a low regeneration temperature (25 °C). In addition, studies 113 done in Materials of Institute Lavoisier (MIL) frameworks showed even greater values of 114 water adsorption  $(1.06-1.7 \text{ cm}^3 \text{ g}^{-1})$  and large pore volume  $(1.4-1.58 \text{ cm}^3 \text{ g}^{-1})^{53-55}$  with low 115 regeneration temperature (30 °C), and rapid-response rates of 40 % weight loss after only 30 116 minutes at 30 °C <sup>55</sup>. It is expected that desirable properties in an optimal desiccant material 117 are therefore not only its water adsorption capacity (and isotherm type), but also their rate of 118 sorption/desorption (kinetic stability) at a defined regeneration temperature. 119

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All of these desiccants have currently only been analysed in terms of full adsorption capacity (*i.e.*, independent of isotherm shape) with the intention of producing large pore volumes, and have not assessed their suitability for specific applications. In order to do this an

understanding of the isotherm shape at particular  $p_v/p_o$  ranges is required, as well as their 124 transient response to changes in ambient vapour pressure,  $\Delta p_{\nu}$ . If understood, advanced 125 desiccant materials could then be used to partially or even fully regulate the fluctuations of 126 127 RH and to allow reduction in energy penalties when tailoring and optimizing their isotherm shape for regulating 'specific' psychrometric requirements of closed environments. Therefore, 128 the aim of this study was to apply a numerical hygrothermal approach (from our previous 129 work <sup>56</sup>) to determine the ideal (and unknown) water vapour isotherm to inform the design of 130 optimised mesoporous desiccants. The isotherms were then experimentally re-created through 131 synthesis and modulation of well-known mesoporous material classes (MCM-41 and SBA 132 133 15). The material's suitability for regulating indoor RH, in terms of dehumidification load reductions (enthalpy), were assessed after different moisture loads had been introduced. They 134 were then assessed to enable the relative effect on energy efficiency when assisting a mixed 135 mode air conditioning (AC) system. The sorption/desorption kinetic response of selected 136 traditional and advanced desiccants will be then assessed experimentally. 137

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#### 139 2 Experimental section

### 140 2.1 Desiccant Candidates

A set of ten candidate desiccants (see Figure 1) were selected as part of a representative group 141 of mesoporous materials, as defined by IUPAC <sup>57</sup>. The aim was to assess their capacity for 142 buffering humidity ratio fluctuations (and hence indoor air enthalpy) within specific closed 143 environment scenarios. In recent work a principle was established which defined the available 144 moisture capacity,  $\Delta w$  for RH buffering within a specific operating range <sup>56</sup>. Therefore, the 145 region of interest among the isotherm candidates was defined between point b seen in Figure 146 2a (1<sup>st</sup> inflection point; the onset of multilayer physisorption) and point c (2<sup>nd</sup> inflection point; 147 the percolation threshold). The capillary transport coefficients ( $D_{ws}$  and  $D_{ww}$ ) were calculated 148

from each isotherm  ${}^{58}$ , and the water adsorption coefficient, *A* was assumed constant for all materials based on assumptions established from previous work  ${}^{38}$ .

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152 2.2 Synthesis and characterization

Synthesis and the textural properties of the pure silica MCM-41@165 were obtained using 153 same experimental description of our previous work <sup>56</sup>. Synthesis of SBA-15 (6 nm pore size) 154 was prepared in a similar manner to Zhao *et al.* <sup>59</sup> with the same batch of SBA-15 used 155 throughout the study. Pluronic 123 (P123, triblock copolymer (EO)20(PO)70(EO)20, 80.0 g) 156 was dissolved in aqueous hydrochloric acid (1.33M, 3000 ml) with vigorous stirring at 35 °C. 157 After complete dissolution tetraethoxyorthosilicate (TEOS, 182 ml) was added dropwise and 158 the mixture stirred at 35 °C for 24 h. The solution was then statically heated at 85 °C for 48 h. 159 The resulting white solid was dried in an oven at 105 °C for 2 h before heating in a furnace to 160 600 °C for 6 h to obtain a white solid powder. Synthesis of SBA-15 (10 nm pore size) was 161 prepared in a similar manner to Zhao et al. <sup>59</sup> with the same batch of SBA-15 used throughout 162 the study. Pluronic 123 (P123, triblock copolymer (EO)20(PO)70(EO)20, 4.0 g) was 163 dissolved in aqueous hydrochloric acid (1.33 M, 75 ml) and mesitylene (2 ml) with vigorous 164 stirring at 35 °C. After complete dissolution tetraethoxyorthosilicate (TEOS, 4.55 ml) was 165 added dropwise and the mixture stirred at 35 °C for 24 h. The solution was then placed in a 166 Teflon lined autoclave and heated at 120 °C for 24 h. The resulting white solid was recovered 167 and then in an oven at 105 °C for 2 h before heating in a furnace at 600 °C for 6 h to obtain a 168 white solid powder. Textural properties (see Table 1) were determined from N<sub>2</sub> sorption 169 analysis at -196 °C using a Micromeritics ASAP 2020 analyser. Samples (approximately 0.2 170 171 g) were outgassed prior to analysis under high vacuum at 120 °C for 15 h. Specific Surface areas (SSA) was calculated using the standard Brunauer-Emmett-Teller (SBET) method, 172 micropore volume (V<sub>m</sub>) by the t-plot method with Harkins and Jura correction for thickness 173 174 curve correction. Average pore diameter  $(D_p)$  was determined by the Barret–Joyner–Halenda

(BJH) method by applying a cylindrical pore model using Kelvin equation with Harkins and
Jura for thickness curve correction. Nitrogen adsorption at relative pressure of 0.99 was taken
as total pore volume (V<sub>p</sub>).

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179 2.3 Dynamic Vapour Sorption (DVS)

180 A gravimetric dynamic vapour sorption (DVS) test was performed to determine the water 181 vapour isotherms for the candidate materials (MCM-41@165, SBA-15 6 nm and SBA-15 10 182 nm). A DVS Advantage-2 series (Surface Measurements Systems Ltd., London, England) with incorporated ultra-microbalance (1µg sensitivity) and fully automated weighing/purging 183 184 was used to independently control  $T_{db}$  and RH to an accuracy of  $\pm 0.5$  K and  $\pm 1.5$  %. respectively (vapour pressure accuracy  $\pm 1.5 \% p_v/p_0$ ). Advantage Control Software (ACS) 185 was used to automatically record the data in 1 min time-steps. Before analysis, the samples 186 were dried under vacuum at 120 °C for 12 h to ensure removal of residual moisture. In order 187 to ensure stability for baseline measurements accuracy, clean quartz sample and reference pan 188 were left to fully equilibrate at 23 °C at constant nitrogen flow (10 sccm 1 bar) for 1 h prior to 189 the weighing and tared. Dry samples (approximately 0.2 g) were weighed into the sample pan 190 and allowed to equilibrate again under the carrier gas for 1 h. After stability, the sample 191 192 weight was recorded (initial mass  $M_0$ ) and reference weight was set to zero with no counter weight. To ensure w = 0, a pre-heating sequence was set from to 23 to 150 °C (2 °C/min 193 ramp rate) for 1 h prior to vapour dosing. A targeted relative vapour pressure from  $\varphi = 0.05$ 194 to 1 ( $\phi = 0.05$  intervals) was programmed to sequentially increase in a 21-steps full cycle 195 (sorption/desorption), where a change in mass step (0.001 %/min) was set until EMC was 196 achieved. This was performed at  $T_{db}$  23 °C and to a constant water vapour flow rate of 100 197 sccm. 198

200 Three sets of experiments to assess the water vapour adsorption/desorption kinetics were

201 performed. For this, a set of RH limits were defined (20–30 %; 30–50 %; and 50–70 %) and

202 the kinetics of the moisture sorption began to sequentially fluctuate in an

adsorption/desorption cycle. Each cycle comprised 2 h of moisture sorption (*i.e.*, 30–50 %

RH) followed by 2 h of moisture desorption (*i.e.*, 50–30 % RH) for a total period of 12 h (3

205 complete cycles). Prior to testing, all samples were outgassed under vacuum at 120 °C for 12

206 h and the pre-heating sequence was applied as described above. To ensure that EMC was

achieved for the  $w_{20}$ ,  $w_{30}$  and  $w_{50}$  set points, all samples were pre-conditioned within the DVS

208 chamber before the cycles started. Taring and equilibration of the microbalance steps were

done as previously described, where all samples ( $\sim 0.02$  g) were loaded to define the initial

mass ( $M_0$ ) in the DVS. This was performed at  $T_{db}$  23 °C at a constant water vapour flow rate

of 100 sccm, as with previous experiments.

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# 213 2.4 Hygrothermal numerical model

A number of hygrothermal models packages have been validated over the past 15 + years against experimental data including (but not limited to) DELPHIN, HAM, MATCH and WUFI, as part of the Annex 41 project <sup>60</sup>. To validate numerical model predictions for heat and mass transport/storage physical model are commonly used to simulate the bulk–scale of the hygrothermal behaviour of some functional material properties <sup>61-63</sup>. As a minimum, all hygrothermal numerical models must accurately predict the following bulk–scale coupled phenomena of heat and mass transport/storage <sup>64</sup>:

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222	•	heat storage in dry materials and any absorbed water
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- heat transport by moisture-dependent thermal conduction
- latent heat transfer by vapour diffusion
- moisture storage by vapour sorption and capillary forces

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- moisture transport by vapour diffusion
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• moisture transport by liquid transport (surface diffusion and capillary flow)

For this study, the Wärme und Feuchte Instationär (WUFI) hygrothermal numerical model <sup>65</sup>, 229 WUFI+ v2.5.3 (Fraunhofer, Germany), was used. The governing equation that represents the 230 moisture storage function is seen in the LHS of Eq. 1, which is proportional to the derivative 231 of the water adsorption isotherm  $(\partial w/\partial \varphi)$ . The transport terms on the RHS of Eq. 1 are 232 described by the divergence of liquid and vapour diffusion with phase changes strongly 233 dependent on the moisture fields and fluxes. Whilst the driving potential for vapour flux, 234 which is vapour pressure ( $p_v = \varphi \cdot p_{sat}$ ), is strongly temperature–dependent (saturation pressure 235  $p_{\text{sat}} \sim \exp(T)$ ). The liquid flow, on the other hand, is governed by differences in capillary 236 forces which are assumed to be a function of RH only (*i.e.*, from Kelvin condensation). The 237 storage term seen in LHS of Eq. 2 describes the vapour enthalpy of air flow inside the closed 238 environment, whilst the RHS describes heat transport due to solid conduction and diffusion of 239 heat associated with the specific enthalpy of vapour water. 240

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242 Moisture balance: 
$$\frac{dw}{d\varphi} \cdot \frac{\partial \varphi}{\partial t} = \nabla \cdot \left( D_w \nabla \varphi + \delta_p \nabla (\varphi \cdot p_{sat}) \right)$$
 Eq. 1

243 Energy balance: 
$$\frac{dH}{dT} \cdot \frac{\partial T}{\partial t} = \nabla \cdot (\lambda \nabla T) + h_v \nabla \cdot (\delta_p \nabla (\varphi \cdot p_{sat}))$$
 Eq. 2

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### 245 2.5 Sensitivity analysis

Sensitivity analysis was conducted to assess the water vapour adsorption behaviour (capacity) and water vapour exchange performance for all candidate materials in a closed environment where the initial moisture content was w = 0. The hygrothermal numerical model was setup to sequentially perform water vapour adsorption simulations between the operating limits  $\varphi_{min} =$ 0 and  $\varphi_{max} = 0.95$ . A moisture generation rate of 3 g/h per 24 h period with and an air

251 infiltration rate of 0.35 ACH h<sup>-1</sup> were set. The water vapour exchange rate was then estimated

as the fraction of cumulative moisture load per hour. The closed environment comprised a 252 thermally and hygrically isolated box (where  $V = 1 \text{ m}^3$ ) with the initial boundary conditions 253 (interior/exterior) were set at  $\varphi_i = 0$  and  $\varphi_e = 0.95$  under isothermal conditions,  $T_{db} = 23$  °C. 254 255 The fabric of the envelope comprised a 2 mm thick vapour barrier ( $S_d = 1500$  m) and 240 mm thick vacuum insulation panel ( $\lambda = 0.001$ ). A non-visualized internal component was defined 256 (area/volume ratio =  $0.4048 \text{ m}^2$ , equivalent to ~500 g) comprising 2 mm thickness of 257 candidate material and using the hygrothermal functional properties detailed in Table 2. These 258 were gathered from previous research <sup>38</sup>, where pore geometry and specific surface area were 259 obtained using the BJH and BET method with N<sub>2</sub> physisorption. Validation against high 260 resolution transmission electron microscopy (HRTEM) and small-angle X-ray diffraction 261 (SAXRD) were conducted. Modified transient plane source (MTPS), differential scanning 262 calorimetry (DSC) and dynamic vapour sorption techniques were used to determine the 263 hygrothermal functional properties. The numerical model was solved using (i) increased 264 accuracy and adapted convergence options, (ii) time step 1 min, 'fine' mesh grid spacing, and 265 (iii) calculation accuracy of 0.5 %/0.5 °C for RH and  $T_{db}$ , respectively. 266

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268 2.6 Parametric studies

Parametric hygrothermal simulations were conducted to determine the fluctuation of  $\varphi(t)$  and 269 the candidate material's capacity for humidity buffering in a series of closed environment 270 scenarios. Each material was parametrically assessed against five pairs of operating limits 271  $(\varphi_{min}-\varphi_{max})$ ; *i.e.*, 0.4–0.5; 0.5–0.6; 0.6–0.7; 0.7–0.8; 0.8–0.9. The initial moisture content, w 272 273 was set according to the  $\varphi_{min}$  limit at EMC; *i.e.*,  $w_{40}$ ,  $w_{50}$ ,  $w_{60}$ ,  $w_{70}$ , and  $w_{80}$ . A number of moisture loading cycles, N was set comprising 12 h moisture generation (at a rate of 3 g/h) 274 followed by 12 h of no moisture generation (24 h total), *i.e.*, 22 cycles in total. The air 275 infiltration rate (0.35 ACH h<sup>-1</sup>) and  $\varphi_e$  were constant, with the latter value determined 276 according to the  $\varphi_{min}$  limit (*i.e.*,  $\varphi_e = 0.4, 0.5, 0.6, 0.7, \text{ or } 0.8$ ) and to a constant dry bulb 277

temperature,  $T_{db} = 23$  °C. The non-visualized component thickness and area/volume ratio, 278 fabric components, and parameters for the numerical model were set as described in Section 279 2.4. Expanded numerical simulations were then conducted to quantify the operational energy 280 281 used for dehumidification loads (enthalpy energy) and cooling demand, with the candidate materials assisting a mixed mode AC system. The upper indoor dry bulb temperature set 282 point,  $(T_{dbi.U})$  was set as  $T_{dbi} \ge 25$  °C, and the upper indoor air humidity set point ( $\varphi_{max}$ ) was 283 defined for each pair of operating limits ( $\varphi_i \ge 0.5, 0.6, 0.7, 0.8, 0.9$ ), as previously described. 284 In order to simulate instantaneous control response, 50 kg/h of dehumidification capacity and 285 50 kW cooling capacity were assumed for the AC system. Once the  $\varphi_{max}$  and  $T_{dbi,U}$  set points 286 had been achieved, and if any of the operational limits were exceeded, the ratio of 287 dehumidification or cooling demand could be estimated and the relative effect on energy 288 efficiency calculated. This commonly occurred in cases where the moisture loads were always 289 positive, *i.e.*, where no infiltration of air at lower relative moisture content occurred. For each 290 candidate material, the initial value of w was set according to the  $\varphi_{min}$  limit, *i.e.*, EMC at  $w_{40}$ , 291 292 w50, w60, w70, and w80. A constant air infiltration rate (0.35 ACH h<sup>-1</sup>) was used, and the moisture load was derived from the resultant outdoor air moisture content at  $\varphi_e = 0.95$  at  $T_{db} =$ 293 23 °C. The non-visualized component thickness and area/volume ratio, fabric components, 294 and parameters for the numerical model were set as described in Section 2.4. The initial 295 boundary conditions were, however, set equally for interior operating limits ( $\varphi_{min}-\varphi_{max}$ ), and 296  $\varphi_e = 0.95$  at  $T_{db} = 23$  °C. The numerical model setup was later adjusted to simulate the 297 kinetics of the water vapour uptake by sequentially fluctuating the adsorption/desorption cycle 298 across an arbitrary sixty cycles (N = 60). Each material was parametrically assessed using the 299 300 same RH limits as for the DVS experiments in Section 2.2 (20-30 %; 30-50 %; and 50-70 %), where the initial w was set accordingly (*i.e.*, EMC at  $w_{20}$ ,  $w_{30}$  and  $w_{50}$ ) and  $T_{db} = 23$  °C. A 301 constant ACH =  $20.37 \text{ h}^{-1}$  and area/ volume ratio (0.01) of the non-visualized component 302 were determined. This was to enable scaling between the DVS chamber volume ( $V_{ch}$  = 303

 $0.0002945 \text{ m}^3$ ) and the numerical model ( $V = 1 \text{ m}^3$ ), assuming an infiltration rate of 100 sccm ( $0.006 \text{ m}^3 \text{ h}^{-1}$ ) and material quantity of 0.02 g. This enabled validation of the numerical model against the DVS experiments.

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# 3 Results and discussion

309 3.1 Continuous moisture load sensitivity analysis

Figure 3 clearly shows three stages of water vapour adsorption as w reaches equilibrium 310 moisture content (EMC), where each stage was identified within a specific portion of the 311 312 isotherm as depicted in Figure 2a. The first stage (schematically sketched in Figure 2b) between points a and b represents an early saturation stage of the materials in the first 48 h 313 where a steep  $\varphi$  gradient was found. In the cases of MIL-101 <sup>66</sup>, PS-C16-R <sup>67</sup> and SBA-15, the 314 higher  $\Delta w$  gradient during adsorption between  $\varphi = 0$  to *ca*. 0.4, and desorption between 0 to 315 ca. 0.9 resulted in a less pronounced curve and a small but significant delay when achieving 316 the  $1^{st}$  inflection point b. A second stage was a sub-plateau between point b and c where a 317 characteristic stabilized  $\varphi_i$  corresponded to the highest  $\Delta w$  at the 2<sup>nd</sup> inflection point c, *i.e.*, the 318 percolation threshold. The slope of the sub-plateau indicates its dependency upon the 319 320 steepness of the w:  $p_v/p_0$  gradient corresponding to maximum pore volume. Previous work has demonstrated that the positive correlation between the moisture storage capacity ( $\Delta w$ ) and the 321 optimum  $\varphi$  operating range for a given isotherm can also be identified <sup>56</sup>. Almost all samples 322 exhibited the sub-plateau segment in their isotherms, which started from point b (onset of 323 multilayer physisorption between  $\varphi = c. 0.35-0.90$ ), with the exception of the Aerogel <sup>68</sup>. This 324 325 suggests a prolonged capillary conduction stage. In the third stage between points c and d, a rapid increase in  $\varphi_i$  was observed with noticeable variation towards EMC<sub>100</sub>, especially in 326 materials having type V isotherms (e.g., KIT-1<sup>69</sup>). The progressive reduction in pore volume 327 strongly affected the time taken to reach inflection point d towards EMC<sub>100</sub>, as a consequence 328 of the low  $\Delta w$  on that portion of the isotherm. Evidence of this was found for MCM-41@180 329

<sup>56</sup>, which was the first to achieve EMC<sub>100</sub> (112 h) and has the lowest pore volume (0.53 g cm<sup>-</sup> <sup>3</sup>). However, it was apparent that by having a greater  $\Delta w$  between points *c* and *d*, the elapsed time before reaching EMC<sub>100</sub> was significantly increased. Evidence of this was found when comparing KIT-1 (152 h) and FSM-16 (168 h) <sup>69</sup>, both with similar  $\Delta w$  at point *c* (460 kg m<sup>-</sup> <sup>3</sup>), but with the latter having a steep *w*:  $p_v/p_0$  gradient resulting in a higher  $\Delta w$  above  $\varphi = 0.7$ (~44 kg m<sup>-3</sup>).

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337 3.2 Water vapour exchange performance

Figure 4 shows the cumulative water vapour uptake for all candidate materials. The same 338 correlations were observed in Section 3.1 for stage 1–3 water vapour adsorption and the 339 corresponding portions of the isotherm (see Figure 2). It appears that in the first stage (points 340 a and b) there is an 'apparent' high initial water vapour uptake (relative to w = 0) followed by 341 a visible depression caused by early saturation. This applied to MCM-41, KIT-1, FSM-16, 342 343 and Aerogel. Note that in the cases of MIL-101, PS-C16-R and SBA-15 the cumulative water vapour adsorption was clearly higher (between 84 % to 99 %) and controlled by the higher 344 initial  $\Delta w$  gradient between points a and b (see Figure 2a). After this stage, a sub-plateau was 345 observed in the second stage (points b and c, with the highest  $\Delta w$  gradient), which 346 corresponded to maximum pore volume. Here some materials (MIL-101, PS-C16-R and 347 SBA-15) reached 100 % of the cumulative water vapour uptake. Once point c of the isotherm 348 was reached the moisture content (with respect to pore volume) was sufficiently high for 349 performance to begin to decay after successive internal moisture load cycles. This indicates 350 351 moisture content dependency and reduced water vapour adsorption ability prior to reaching  $w_{100}$  from point c of the isotherm. Figure 5a shows a positive linear correlation between the 352 moisture storage capacity, w and the water vapour being exchanged (against time) with the 353 absorbent. However, Figure 5b shows that the relationship between w and the time taken to 354 exceed the  $\varphi_{max}$  limit ( $\varphi = 0.95$ ) correlates more strongly with type V isotherms (R<sup>2</sup> = 0.99) 355

compared with type III isotherms ( $R^2 = 0.77$ ). This could partially be explained by the observation that the process of water vapour adsorption was triggered early by a capillary condensation stage on Type III isotherms. Therefore, SBA-15 10 nm and Aerogel material do not fit trend line A since their capillary potential and capacity for water vapour adsorption are already lowered.

361

## 362 3.3 Water vapour buffering simulations

Figure 6a–b shows the RH buffering results for repeated long-term loop cycling (at N = 12) of 363 all material candidates. The operating range ( $\varphi_{min}-\varphi_{max}$ ) of each test was 0.4–0.5 and 0.7–0.8, 364 respectively. The absence of water vapour buffering can be seen followed by an early decay 365 process which correlates to the first stage of water vapour adsorption between points a and b 366 of the isotherm (Figure 2a). This sudden decay relates to the smaller  $\Delta w$  gradient seen in 367 Figure 7a, causing the material to fail after N number of cycles when the fluctuation of  $\varphi(t)$ 368 exceeds the  $\varphi_{max}$  limit (Figure 7b). From Figure 7b it is apparent that the progressive decay 369 until material failure corresponds with the largest proportion of water vapour adsorption (see 370 Figure 7a) in each operating range. The most suitable candidate materials for each range can 371 372 be identified as that with the longest time before exceeding the  $\varphi_{max}$  limit. This suggests that the sub-plateau segment (see y-axis displacement in Figure 3) is sensitive both to x-axis 373 adjustments in the sorption isotherm (*i.e.*, pore size distribution) and to the value of  $\Delta w$  (w: 374  $p_{\nu}/p_0$  gradient) between  $\varphi_{i,L}-\varphi_{i,U}$  limits. 375

376

377 3.4 Air dehumidification and energy consumption

Figures 8a–c show the ratio of dehumidification loads removed by the materials once they

achieve the  $\varphi_{max}$  set point on the mixed mode AC system (repeated for ranges between: 40–

50; 60–70; and 70–80 % RH). The steepness of the curve was found to be dependent on the

time required to achieve the corresponding EMC (*e.g.*, EMC<sub>50</sub> as seen in Figure 8a). This

supports the idea that total control (100%) from the AC system corresponds to the lower  $\Delta w$ 382 gradient, whereas a plateau (at 0 %) means that the indoor moisture load was fully regulated 383 by the desiccant by having a higher  $\Delta w$  gradient. In all cases, it can be recognized that a 384 385 higher elapsed time before the inflection of the curve start there is a corresponding material that fits within the optimum operating ranges; for example, MIL-101 and PS-C16-R ( $\varphi = 0.4$ -386 0.5), MCM-41@150<sup>56</sup>, KIT-1, and FSM-16 ( $\varphi = 0.6-0.7$ ), and all MCM-41 types ( $\varphi = 0.7-$ 387 0.8). Once the inflection point starts (above 0 %; see red circles in Figure 8a) in the sub-388 plateau region, the adsorbent progressively has lower potential to fully regulate  $\varphi_i$  and air 389 dehumidification starts to operate. 390 Figure 9 shows a significant difference between the resultant indoor RH(t) and the material's 391 RH(t) in all simulations. Since the adsorbent appears not to have achieved EMC (*i.e.*, EMC<sub>50-</sub> 392 <sub>70-80</sub>) a distinctive offset between both curves was observed. This suggests that the material 393

may continue adsorbing water vapour whilst the  $\varphi_{min}$  set point on the AC is still

dehumidifying the indoor air, and hence working in true mixed mode.

396

Figure 10 shows a positive linear correlation between the moisture sorption capacity ( $\Delta w$ 397 gradient) and the latent heat used for air dehumidification for each specific operational range. 398 399 This satisfactorily proves that passive regulation of RH fluctuations in a specific closed environment can fully be controlled by materials where a higher  $\Delta w$  gradient is found whilst 400 consuming less energy. This was previously evidenced in Figures 8a-c, where reduction of 401 the dehumidification loads removed by the HVAC system was strongly related to a higher  $\Delta w$ 402 gradient corresponding to materials that fit within the optimum operating ranges. Consistent 403 404 agreement was also observed between the decay (up to material failure) and the latent heat of dehumidification, wherein the most suitable candidate materials can be identified in Figure 405 7b. A significant increment in the resultant indoor air temperature was observed in some cases 406 407 (e.g., MIL-101 and PS-C16-R), most likely due to heat of adsorption. As a result, an extra

cooling load was added to both MIL-101 and PS-C16-R (i.e., 0.22 and 0.16 KW respectively) 408 in order to maintain isothermal conditions ( $T_{dbi} = 23$  °C). Figure 11 shows the total energy 409 used for air dehumidification (latent heat) and the energy used for cooling purposes (Figure 410 11a), and no correlation trend ( $R^2 = 0.2706$ ) between total moisture storage, w and total latent 411 heat for air dehumidification (Figure 11b). In the latter case, it can be seen that having higher 412 413 w (e.g. Aerogel, MIL-101, PS-C16-R and SBA-15 10 nm) does not result in lowering the 414 energy used for dehumidification purposes when assuming total pore volume capacity. 415 Evidence of this can be seen when comparing the total latent heat used for type V and type III isotherms; *e.g.*, MCM-41@180 (1.51 kW and w = 369.4 kg m<sup>-3</sup>) and Aerogel (1.65 kW and w 416 417 = 754.8 kg m<sup>-3</sup>), with the latter having  $w \sim 2$  times higher compared to MCM-41@180. These results provide evidence for the premise in which adsorbents are expected to fully operate 418 under total adsorption capacity, independent of isotherm shape, rather than for specific 419 applications and  $p_{\nu}/p_0$  scenarios. 420

421

422 3.5 Experimental work and model validation

Figure 12 compares the experimental DVS and numerical simulations (water vapour

424 sorption/desorption kinetics) for the three newly synthesised materials (MCM-41@165, SBA-

425 15 6 nm and SBA-15 10 nm). The upper and lower % RH buffering limits were defined as

426 20–30 % RH; 30–50 % RH; and 50–70 % RH in Figure 12. Experimentally, it appears that for

all materials EMC (red dotted line in Figure 12) was achieved at N = 3 for cases where the  $\Delta w$ 

428 gradient was sufficiently small (a, b, d, e, and g) with respect to the relevant portion of each

sorption isotherm (see Figure 1). Additionally, it appears that the kinetics for

430 sorption/desorption within the % RH buffering limits was highly sensitive to the  $\Delta w$  gradient.

431 This corresponded to a rapid response (steep adsorption/desorption curve) for the smallest  $\Delta w$ 

432 gradient. Evidence of this was found in MCM-41@165 operating at 20–30 and 30–50 % RH

433 (see red boxes in Table 3). In all other cases it was evidenced by a less steep

sorption/desorption curve and corresponding to those with a higher  $\Delta w$  gradient. To some 434 extent, the numerical predictions seen in Figure 12 (a, b, d, e, and g) appear to be slightly 435 underestimated (solid black line) towards the time in achieving EMC, especially in those 436 437 between  $\Delta w_{20-50}$  (see Table 3). This is normally found in numerical models due to the lack of a dynamic input for w, with respect to the time-response for water vapour adsorption at partial 438 439 EMC; *i.e.*, w inputs are assumed to be at full EMC for single sorption isotherms. Further analysis confirmed that the average difference (%) between experimental and numerical w 440 441 data for all sorption cycles was significantly attributed to this; e.g., MCM-41@165 with 43.4% within the  $\Delta w_{30-50}$  cycle seen in Table 4. In the case of those having greater w between 442 443 the upper and lower % RH buffering limits (c, f, h, and i from Figure 12), it is suggested that for both experimental and numerical data, their capacity for water vapour adsorption is 444 significantly beyond the required time period for achieving EMC (see Table 3). Besides, for 445 all candidates seen in red circles (Figure 12), there appears to be a combination of time-446 response together with a hysteresis phenomenon. The latter, can be recognized by the highest 447 448 average (%) values at the end of the third cycle shown in Table 4 (see a, c, d, and g). In both adsorption and desorption cycles, this could be largely due to the presence of scanning curves 449 within the hysteresis loop. However, the corresponding loops at this stage are unidentified. 450 451 Conversely, there is close agreement (full sorption/desorption reversibility) when a smaller  $\Delta w$  gradient is found; for example, SBA-15 10 nm operating in the 50–70 % RH ranges (see 452 Figure 12i). This could be explained because SBA-15 10 nm showed the lowest average 453 difference between experimental and numerical w data (see Table 4 - 7.6%), which indicates 454 a high degree of sorption/desorption reversibility (and small hysteresis -4.6%). On the other 455 456 hand, numerical data of the final EMC were accurately predicted (refer to Section 3.1), suggesting that there is some level of variance between short and long periods for water 457 vapour adsorption/desorption kinetics. This perhaps indicates a need for further research in 458 459 which the concept of time-response for water vapour adsorption, hysteresis loops and the

associated scanning curves can be fully implemented by the current state-of-the-art
hygrothermal numerical models.

462

### 463 **4 Conclusions**

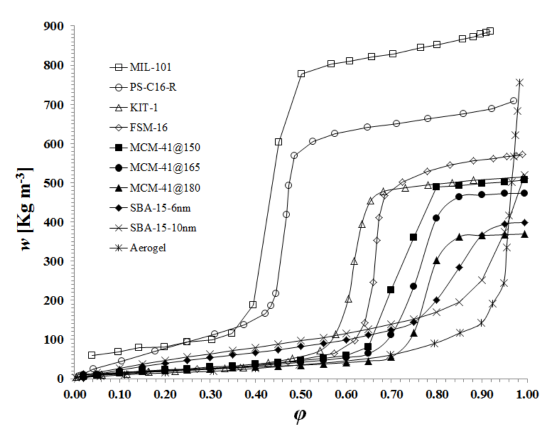
Three new mesoporous desiccant materials (MCM-41@165, SBA-15 6 nm and SBA-15 10 464 nm) were successfully synthesised with pore size modulation, in order to experimentally 465 recreate the ideal isotherms determined (for differing applications) by the hygrothermal model 466 optimization technique previously developed by the authors. A series of numerical 467 simulations were performed to assess the desiccants' capacity for regulating indoor RH and 468 the corresponding impact on energy efficiency when assisting a mixed mode AC system. The 469 simulated results clearly showed three stages of water vapour adsorption that strongly related 470 to the  $\Delta w$  gradient showing a positive linear correlation with the latent heat used for air 471 dehumidification. The results were also consistent with the predictions for progressive decay 472 and the point of failure, indicating the suitability of the materials for the  $p_{y}/p_{0}$  scenarios for 473 474 which they had been designed. Increasing w did not result in lowering the energy used for 475 dehumidification purposes, especially for the Type III isotherm materials, and found no correlation ( $R^2 = 0.2706$ ) between total moisture storage, w and total latent heat for air 476 477 dehumidification. Experimental DVS and numerical simulation for water vapour sorption/desorption kinetics were conducted for the three new materials, in which the latter 478 appeared to be slightly underestimated. This was consistent for most desiccants shown in 479 Figure 12 and Table 4, where the lack of a dynamic input for w (time-response for water 480 481 vapour adsorption), caused significant differences between both results. Additionally, Table 4 482 showed (see a, c, d, and g) that the major underestimations were largely attributed to a combination of adsorption kinetics (time-response) and hysteresis. This, for the case of 483 MCM-41@165, was found to account for 52.4% (average) of the total w values at the third 484 485 RH buffering cycle. Conversely, it was found that the numerical model accurately predicted

486	EMC	and found very good agreement with the experimental data taken from the water vapour
487	isothe	erm. Further research is therefore needed for coupling both adsorption kinetics ( <i>i.e.</i> ,
488	adsor	ption time-response) and realistic scanning curve estimates within the hysteresis loop.
489		
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495	6 F	References
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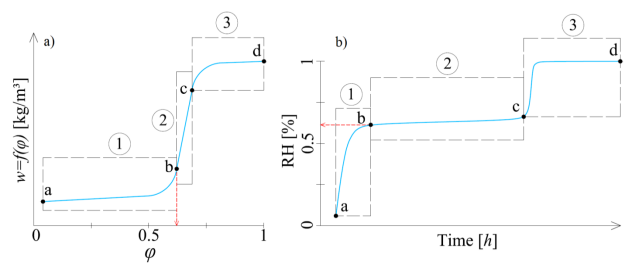
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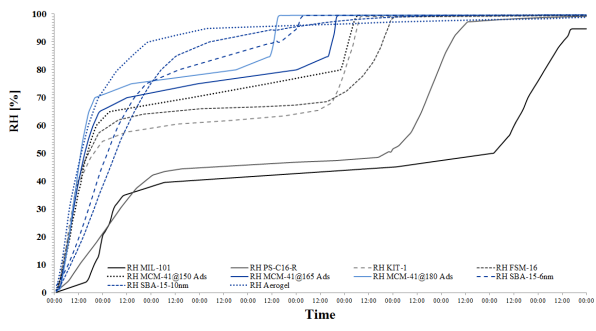


**Figure 1** – Water vapour sorption isotherms for all desiccant candidates. References are provided for the following desiccants candidates: MIL-101<sup>66</sup>, PS-C16-R<sup>67</sup>, KIT-1 and FSM-16<sup>69</sup>, MCM-41@150 and 180<sup>56</sup>, Aerogel<sup>68</sup>. The three water vapour isotherms from the new materials made (MCM-41@165, SBA-15 6 nm and 10 nm) are also provided.

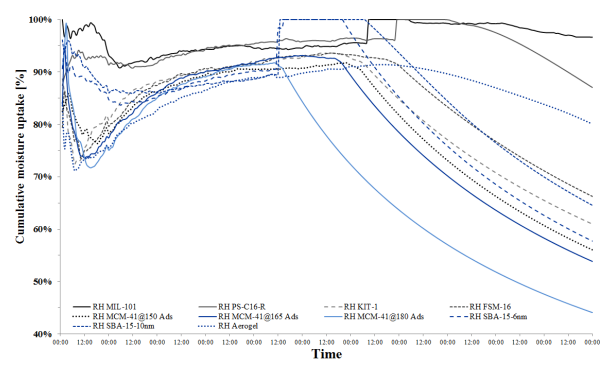


**Figure 2** – (a) Schematic representation of a water vapour sorption isotherm portions at different EMC stages a-b, b-c and c-d. (b) Corresponding water vapour sorption kinetics at different EMC stages a-b, b-c and c-d that shows EMC in the segments 1, 2 and 3.

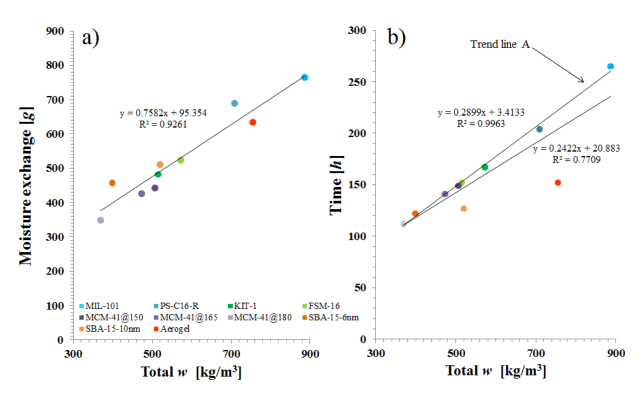




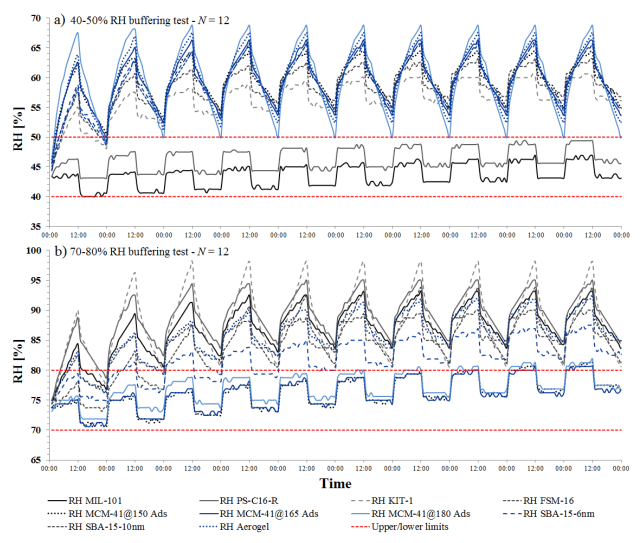
**Figure 3** – Water vapour sorption kinetic stages for all desiccant candidates when achieving EMC<sub>100</sub>. Initial w = 0 for all desiccant materials and the moisture load applied is the moisture content of the infiltrated air at ACH = 0.35<sup>-1</sup>, where  $\varphi_e = 0.95$ .



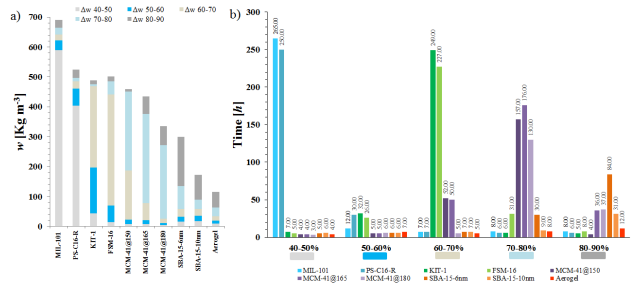
**Figure 4** – Cumulative water vapour uptake for all desiccant candidates. Initial w = 0 for all desiccant materials and the moisture load applied is the moisture content of the infiltrated air at ACH = 0.35<sup>-1</sup>, where  $\varphi_e = 0.95$ .



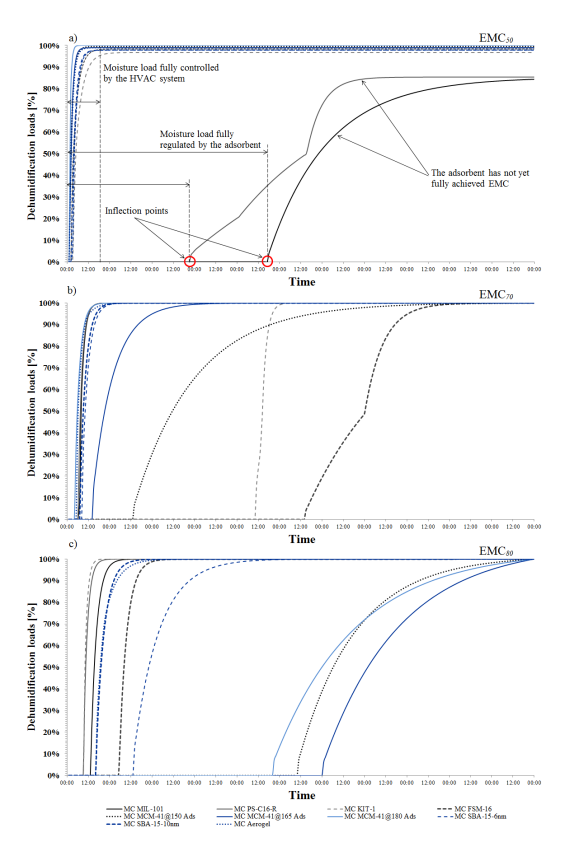
**Figure 5** – (a) Linear correlation between the moisture storage capacity, w and the water vapour being exchanged. (b) Relationship between total moisture storage capacity and time taken to achieve EMC<sub>100</sub>.



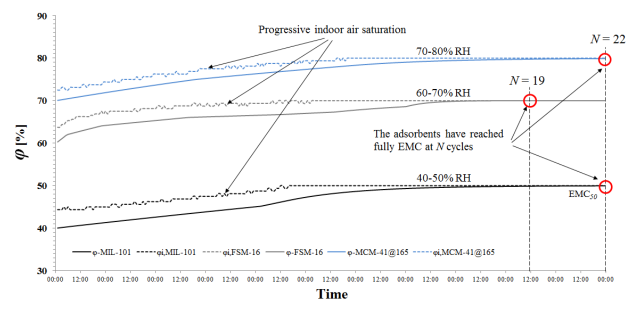
**Figure 6** – RH buffering results for all desiccant candidates, where the number of cycles, N = 12. (a) Shows the simulations results for RH buffering between 40–50%, and (b) 70–80% operating ranges. The red-dotted lines correspond to the maximum and minimum RH limits.



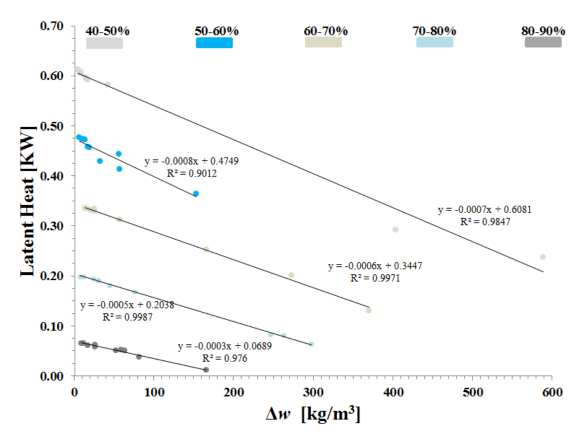
**Figure 7** – (a) Water vapour uptake capacity at different  $\Delta w$  for all desiccant candidates ( $\Delta w = 40$ -50; 50-60; 60-70; 70-80; and 80-90). (b) Time taken for all desiccant materials when the fluctuation of  $\varphi(t)$  exceeds the  $\varphi_{max}$  limit; where  $\Delta w = 40$ -50; 50-60; 60-70; 70-80; and 80-90. This also correspond to the threshold when the materials reaches failure at the number of cycles, N = 60.



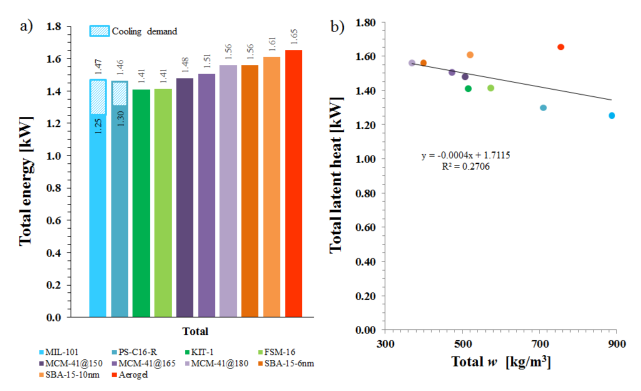
**Figure 8** – Dehumidification ratio performed by the HVAC system and the desiccant candidates according to the operating ranges (a) 40–50; (b) 60–70; and (c) 70–80 % RH. Here, 0% indicates that the desiccant candidate is fully regulating indoor RH. Above 0% indicates that the HVAC system start to working in truly mixed mode. The time taken for reaching EMC at the corresponding operating ranges are solely dependent upon the  $\Delta w$  gradient.



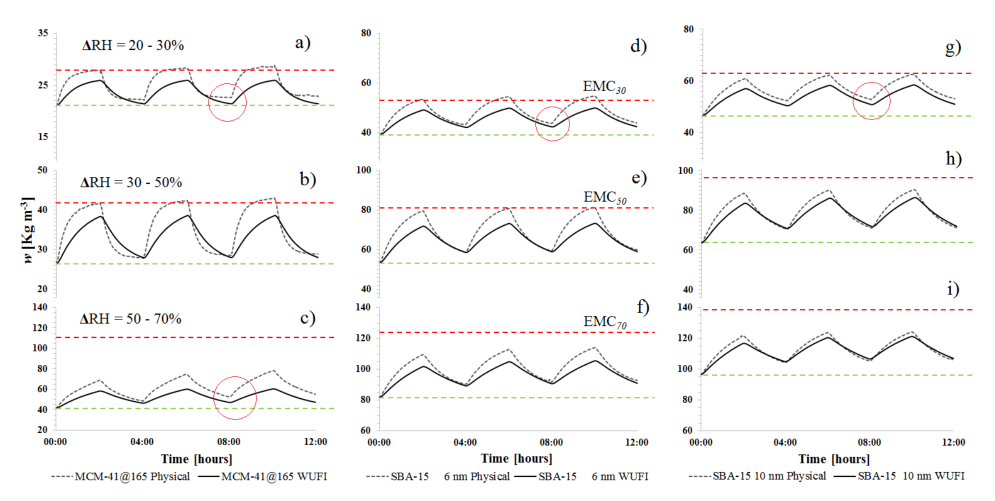
**Figure 9** – Comparison between the resultant indoor RH(*t*) and the material's RH(*t*) at different operating ranges for desiccant materials MIL-101, FSM-16 and MCM-41@165. The offset dotted-lines represent the progressive indoor air saturation towards equilibrium with the desiccant at the respective ACH <sup>-1</sup>. The total number of cycles, N = 22 and the red circles represents when the desiccants have been reached EMC at the corresponding operating RH ranges.



**Figure 10** – Correlation between moisture sorption capacity and the latent heat used for air dehumidification from the HVAC system at the corresponding operating ranges (40-50%; 50-60%; 60-70%; 70-80%; 80-90%). Each trend line correspond to the aggrupation of all desiccant candidates when performing the air dehumidification through the HVAC system.



**Figure 11** – (a) Total energy used for air dehumidification for all desiccant candidates. In the case of materials MIL-101 and PS-C16-R, a cooling demand is represented by the hatched box above the values. (b) Non-linear correlation between total moisture storage, *w* and total latent heat for air dehumidification. Type III isotherms can clearly be recognized as the precursors of this non-linear trend as their total moisture content, *w* do not correlate with the total energy used for air dehumidification and cooling loads, resulting in  $R^2 = 0.2706$ .



**Figure 12** – Water vapour sorption/desorption kinetics comparison between experimental DVS and numerical simulations. The results are deemed to be read horizontally, where A) indicates the operating range  $\Delta RH = 20-30\%$ ; B)  $\Delta RH = 30-50\%$ ; and C)  $\Delta RH = 50-70\%$ . The red-dotted lines indicate achieving the corresponding EMC, whereas the green-dotted lines indicate the base line for the initial *w*.

Sample	$S_{BET}\!/m^2g^{\text{-}1}$	$V_m/cm^3g^{-1}$	$V_p/cm^3g^{-1}$	D <sub>p</sub> /nm
SBA-15 6 nm	876	0.17	0.91	5.7
SBA-15 10 nm	757	0.12	1.26	10.2

**Table 1** – Textural properties summary for SBA-15 desiccant candidates.

Material	п	$\phi_{pore}$	$\rho_b$	$C_p$	λ	Α	μ
	(m <sup>3</sup> /m <sup>3</sup> )	(nm)	$(kg/m^3)$	(J/kg K)	(W/m.K)	$(kg/m^2 s^{0.5})$	(-)
MS-8.3*	0.76	8.3	618	1691	0.05	0.550	10.52
Table 2 – H	ygrothermal fi	unctional	properties	for meso	porous silic	a. *Values tal	ken fron

reference  $^{38}$ .

Desiccant materials	Moisture storage at $\Delta w$			Time in achieving EMC			
	$\Delta w_{20-30}$	Δ <i>w</i> 30-50	$\Delta w$ 50-70	Δw20-30	Δ <i>w</i> 30-50	$\Delta w$ 50-70	
		$(kg/m^3)$		(~ h)			
MCM-41@165	5.4	15.1	69.8	3	5.5	22	
SBA-15 6nm	14.2	28.3	42.5	5	9.5	11	
SBA-15 10nm	16.8	33.1	41.8	6	11	11	

**Table 3** – Moisture storage capacity at different  $\Delta w$  for all new desiccant materials. The time in achieving EMC is also presented with an approximate time (h) for each desiccant at the corresponding ranges:  $\Delta w_{20-30}$ ;  $\Delta w_{30-50}$ ; and  $\Delta w_{50-70}$ . These time values represent numerical predictions based on *w* inputs that are assumed to be at full EMC.

Mesoporous desiccant materials									
	MCM-41@165			SBA-15 6 nm			SBA-15 10 nm		
Cycles	Δw20-30 Δw30-50 Δw50-70		$\Delta w_{20-30} \ \Delta w_{30-50} \ \Delta w_{50-70}$		Δw20-30	$\Delta w_{20-30} \Delta w_{30-50} \Delta w_{30-50}$			
	a	b	C	d	e	f	g	h	i
Ads. cycle 1	44.4%	43.3%	48.4%	44.4%	43.6%	39.8%	40.0%	35.2%	27.8%
Des. cycle 1	11.3%	-56.6%	28.2%	16.9%	6.5%	13.0%	24.9%	1.7%	3.0%
Ads. cycle 2	48.8%	40.5%	45.9%	38.2%	32.2%	28.6%	33.1%	18.5%	11.5%
Des. cycle 2	29.4%	-27.9%	41.8%	21.3%	7.3%	12.0%	23.2%	-3.7%	-4.2%
Ads. cycle 3	51.8%	43.4%	52.4%	38.1%	33.4%	29.0%	32.7%	16.6%	7.6%
Des. cycle 3	36.3%	-15.0%	49.2%	23.9%	11.0%	14.8%	24.1%	-3.7%	-4.6%

**Table 4** – Average difference (%) between experimental and numerical *w* data for all new mesoporous desiccant materials, when modelling for each independent RH buffering scenario 20–30 % RH; 30–50 % RH; and 50–70 % RH. The *w* differences are provided for the following ranges:  $\Delta w_{20-30}$ ;  $\Delta w_{30-50}$ ; and  $\Delta w_{50-70}$ , denoted with the vowels (a–i) according to Figure 12, and for the three completed adsorption/desorption cycles.