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# Different bed configurations and time ratios: Performance analysis of low-grade heat driven adsorption system for cooling and electricity

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## Abstract

In this study, different multi-bed water adsorption systems have been used to generate cooling and electricity at the same time using 9 different cases including 7 bed configurations and 7 time ratios ( $R = \text{total switching and adsorption time} / \text{the total switching and desorption time}$ ) utilizing advanced adsorption materials such as AQSOA-Z02 and MOF Aluminium-Fumarate additionally to traditional Silica-gel. A MATLAB Simulink program of multi-bed adsorption system for cooling and power generation has been developed to investigate the effect of using different cases on the overall system performance. Results showed that using three-bed configuration with time ratio of ( $R=1/2$ ) produced the highest specific cooling power (SCP) and specific power (SP) for Silica-gel (for all heat source temperature range), Aluminium-Fumarate (for heat source temperature higher than 120 °C) and AQSOA-Z02 (at heat source temperature of 160 °C). Moreover, using two-bed configuration with time ratio of ( $R=1$ ) generates the highest coefficient of performance (COP) for all adsorption materials within the range of heat source temperature used in this study. Results also, showed that maximum COP of 0.64 can be achieved using Silica-gel, while maximum SCP, SP and adsorption power efficiency of 650 W/kg<sub>ads</sub>, 64 W/kg<sub>ads</sub>, 4.6 % can be achieved using AQSOA-Z02.

Keywords: Multi-bed, Time Ratio, Adsorption, Cooling and Electricity, AQSOA-Z02, Aluminium-Fumarate, Silica-gel

## 1. Introduction

Nowadays, searching for alternative energy sources becomes an essential aim because of the large energy consumption around the world, while electricity is still mainly generated by burning fossil fuel which rises CO<sub>2</sub> emissions. In hot countries, refrigeration and air conditioning equipment used in houses and offices consume a large amount of electricity, while millions of people, especially in poor countries still face difficulties to access low-cost and reliable energy sources. Adsorption technology offers the potential of using low-grade heat sources such as solar, geothermal and waste heat from industrial processes to produce cooling and electricity simultaneously. Fossil fuel is still the main source of energy since decades; however, the abundant renewable resources such as low-grade heat from solar energy or waste

35 heat from industrial processes can be converted into useful cooling and electricity using promising clean  
36 energy technologies like absorption, adsorption and Organic Rankine cycle (ORC). Such renewable  
37 technologies can be improved to face the serious concerns of the global warming around the world and  
38 reduce the emissions of the CO<sub>2</sub>. A number of researchers investigated the absorption cooling technology  
39 experimentally [1] and numerically [2], while many researchers investigated means of improving the  
40 adsorption cooling technology using multi-bed [3], multi-stages [4], utilizing a number of working pairs  
41 [5] through modelling [6] and experimental work [7]. Gonzalez-Gil et al [8] experimentally studied a  
42 solar air-conditioning water/lithium bromide absorption system and reported a COP of about 0.6 with a  
43 cooling capacity between 2 and 3.8 kW. Wang [9] used new ideas to integrate heat pipes with adsorption  
44 water chiller achieving average cooling and COP of 10 kW and 0.4 using heat source temperature of 85  
45 °C. Gong et al [10] examined experimentally the cooling effect of lithium chloride and Silica-gel  
46 composite adsorption chiller with methanol. Results showed that, compared to the Silica-gel/water unit,  
47 the SCP and COP of the composite were increased by 16.3% and 24.2%.

48 Generating power (electricity) at high efficiency using low-grade heat sources like solar energy,  
49 geothermal energy and waste heat from industrial processes is still a strategic goal for many researchers.  
50 Kalina cycle and Organic Rankine cycle (ORC) have the capability of generating electricity using such  
51 low-grade heat sources [11, 12]. Le et al. [13] investigated a number of ORC fluids to optimize the  
52 performance of basic and regenerative supercritical ORC utilizing heat source temperature of 150 °C.  
53 Results showed that the maximum efficiencies of the basic and regenerative were 11.6 % and 13.1 %  
54 respectively using R152a, while the maximum power generated was 4.1 kW using R1234ze.

55 In order to achieve both cooling and electricity at the same time, a number of researchers designed and  
56 investigated combined systems for cooling and electricity utilizing low-grade heat sources.  
57 Vijayaraghavan and Goswami [14] studied two configurations to combine an absorber and a turbine in  
58 one system to generate cooling and electricity. Results showed that efficiency can be enhanced by up to  
59 25%, while exergy analysis showed that RUE (resource utilization efficiency) can be improved. Liu and  
60 Zhang [15] have proposed a cogeneration system consisting of ammonia/water Rankine cycle and an  
61 ammonia absorption refrigeration cycle utilizing heat source temperature of around 450 °C to generate  
62 cooling and power simultaneously. Results showed that maximum exergy efficiency of 58% can be

63 achieved and the energy consumption can be reduced by 18.2% compared to conventional separate  
64 systems. Zheng et al. [16] have suggested a combined absorption cooling and power system based on  
65 Kalina cycle's principles. The cycle produced cooling and power at the same time with an overall thermal  
66 and exergy efficiencies of 24.2% and 37.3% respectively. Zhang and Lior [17] investigated an  
67 ammonia/water cogeneration system to produce cooling and power at the same time. The system works in  
68 parallel mode with an ammonia/water Rankine cycle and an ammonia refrigeration cycle. Energy and  
69 exergy efficiencies were assessed and they had the values of 27.7% and 55.7%, respectively using heat  
70 source temperature of 450 °C. Absorption technology can generate cooling and electricity simultaneously,  
71 however, such systems have large size, besides ammonia is a toxic material which may cause serious  
72 health risks. On the other hand, adsorption technology has robust construction, ease of installation, and in  
73 many cases, is considered to be more advantageous than absorption systems. In addition, There are no  
74 opportunities of crystallization, corrosion, risky leaks and the power consumption is negligible [18].

75 Adsorption is promising technology that recently used to generate cooling and electricity at the same time  
76 utilizing low-grade heat sources. This technology has a number of advantages like very few moving parts,  
77 high reliability, capability over a wide range of heat source temperatures with appropriate adsorption  
78 materials, and environmentally friendly refrigerants [19]. Jiang et al. [20] have introduced a resorption  
79 cogeneration system to generate cooling and power simultaneously. Results showed that with using heat  
80 recovery the cycle can achieve electrical efficiency and COP of 9.5–15.8% and 0.416–0.691 respectively.  
81 Wang et al. [21] have presented a novel resorption cogeneration cycle for cooling and electricity  
82 simultaneously based on ammonia adsorption cooling technology. Results showed that utilizing a driving  
83 temperature higher than 100 °C can produce a maximum overall exergy efficiency of 0.9 and a COP of  
84 0.77. Bao et al. [22] have studied an adsorption cogeneration prototype to generate power and cooling  
85 simultaneously. The COP and the exergy efficiency of the cycle are 0.57 and 0.62 respectively. Bao et al.  
86 [23] built chemisorption cogeneration prototype using calcium chloride and activated carbon was to  
87 generate power and cooling simultaneously. Results showed that the system could achieve a minimum  
88 value of 5.4 °C at the evaporator and it could produce 490 W of power. L. Jiong et al. [24] have designed  
89 and studied a resorption cogeneration cycle for cooling and power using a scroll expander with  $\text{MnCl}_2$ -  
90  $\text{CaCl}_2$ - $\text{NH}_3$ . Results showed that maximum power of 300W and cooling of 2kW as well as 91 min of cold

91 storage function can be achieved. Yiji Lu et al [25] improved the ammonia resorption cogeneration cycle  
92 proposed by Wang et al with mass and heat recovery using twelve different working pairs. Results  
93 showed that COP increased by 38% and 35% utilizing  $\text{NiCl}_2$  and  $\text{MnCl}_2$  respectively. The efficiency of  
94 electricity has also been improved from 8% to 12% and the second law efficiency reached 41% using  
95  $\text{BaCl}_2\text{-MnCl}_2$ . Al- Mousawi et al. [26] simulated a two-bed adsorption system to generate cooling and  
96 power simultaneously using AQSOA-Z02/water, MIL101Cr/water and Silica-gel/water using heat source  
97 temperature between 80 and 160 °C. Results showed that, the system can generate SP of 73 W/kg<sub>ads</sub>, and  
98 SCP of 681 W/kg<sub>ads</sub> (using AQSOA-Z02) and maximum efficiency of 67%. Al-Mousawi et al. [27]  
99 designed and modelled a small scale radial inflow turbine with efficiency of 82% to generate electricity of  
100 785 W in addition to cooling from a two-bed adsorption system utilizing AQSOA-Z02/water using heat  
101 source temperature of 160 °C. Al-Mousawi et al. [28] studied integrated adsorption-ORC system to  
102 simultaneously generate cooling and electricity. Results showed that system can achieve efficiency, SP  
103 and SCP of 70%, 208 W/kg<sub>ads</sub> and 616 W/kg<sub>ads</sub> respectively.

104 In adsorption cooling systems, as the cycle time increases the coefficient of performance increases, and  
105 this decreases the specific cooling power (SCP) [29]. Many researchers have studied the performance  
106 optimisation of two-bed adsorption cooling systems [29, 30] and three-bed adsorption cycles [31].  
107 Glaznev and Aristov [32] found experimentally that desorption process is faster than adsorption process  
108 by 2.2 to 3.5. Sapienza et al. [33] experimentally found that the best performance can be achieved with  
109 the adsorption time is 7 times longer than desorption timer using driving temperature of 90 °C.  
110 Zajaczkowski [34] found that in a three-bed adsorption system and for switching time 30 s and adsorption  
111 time 300 s, the desorption/adsorption time ratio is almost 0.6 gives the highest improvement in SCP and  
112 COP. Graf et al. [35] showed that the adsorption and desorption times can affect the COP and SCP and  
113 they found that the maximum values of COP and SCP are 268 W/kg and 0.51 respectively using  
114 adsorption time of 200 s and desorption time of 125 s.

115 None of the previous work studied the effect of using different bed configurations and bed arrangement  
116 either in series or in parallel besides the effect of adsorption/desorption time ratio on the overall  
117 performance of the adsorption system for cooling and electricity. In this paper, 9 different cases  
118 including 7 different configurations and 7 time ratios have been investigated. Different cases are

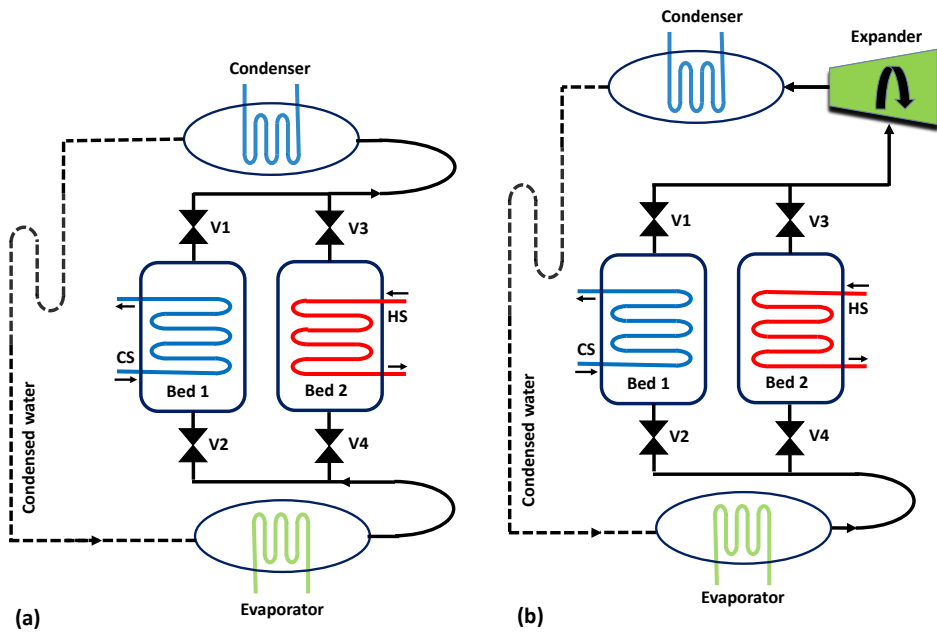
119 compared to the two-bed adsorption system for cooling and electricity in terms of system coefficient of  
120 performance (COP), specific cooling power (SCP), specific power generated (SP) and adsorption power  
121 efficiency. In addition, advanced adsorption pairs of AQSOA-ZO<sub>2</sub>/water and Aluminium-Fumarate  
122 MOF/water have been investigated and compared to Silica-gel/water, while environmentally friendly  
123 fluid of water has been used as adsorption working fluid.

## 124 **2. Two-bed adsorption system for cooling and electricity**

125 Figure 1(a) shows a schematic diagram of a basic cooling adsorption system (BCAS) which contains two  
126 adsorption beds, condenser, and evaporator. Adsorption is an exothermic process, as a result cooling is  
127 needed to cool the adsorber (cold bed) during the adsorption process using cooling source CS in order to  
128 adsorb the refrigerant from the evaporator and produce the cooling capacity. Desorption is an  
129 endothermic process, so heating is needed during the desorption process to release the refrigerant (water  
130 vapour) from the desorber (hot bed) using heating source HS like solar energy or waste heat.  
131 Subsequently, condenser cools the hot refrigerant coming from the desorber to feed the evaporator with  
132 liquid refrigerant that needed to produce cooling continuously.

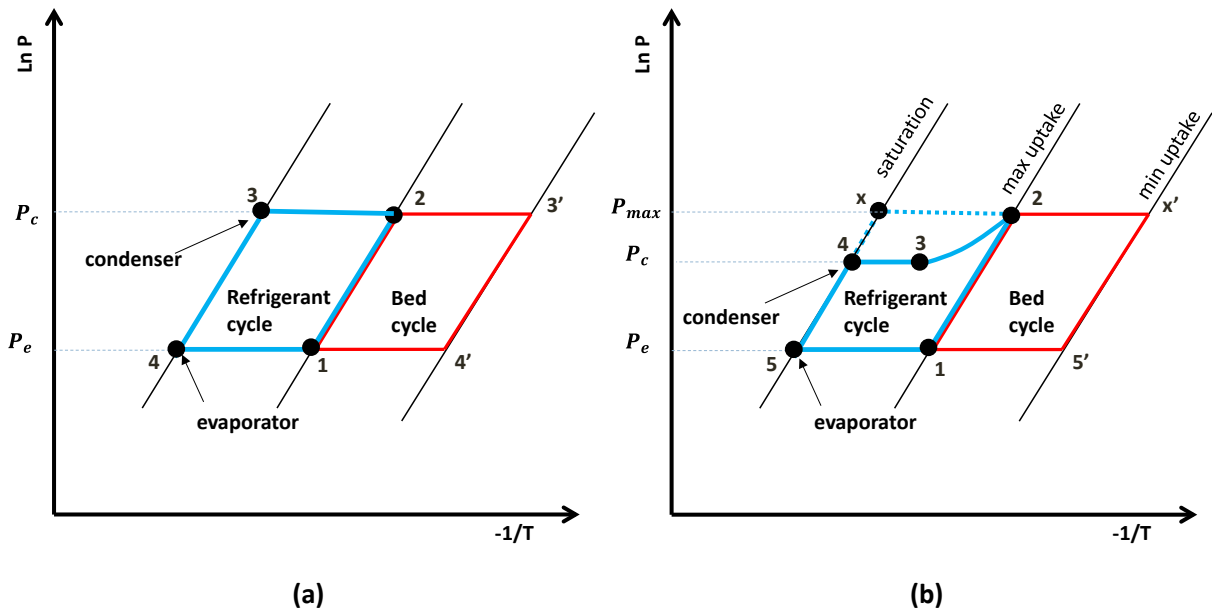
133 Figure 1(b) shows a schematic diagram of the adsorption system for cooling and electricity as described  
134 by a number of literature [22, 26, 27, 36] which consisting of adsorber, desorber, condenser, evaporator  
135 and expander (turbine) located between the hot bed (desorber) and the condenser to extract the kinetics  
136 energy from the refrigerant vapour at high temperature and pressure. Figure 2(a) shows the P-T diagram  
137 of BCAS, process 1-2 is isosteric heating (preheating switching), processes (2-3'/2-3) are isobaric  
138 desorption/condensation, process 3'-4' is isosteric cooling (precooling switching) and finally processes  
139 (4'-1/4-1) are isobaric adsorption/evaporation. Figure 2(b) shows the P-T diagram of the adsorption  
140 system for cooling and electricity and in this system, the pressure of the condenser have to be lower than  
141 the pressure of the hot bed to make the pressure difference required to generate power (electricity) in the  
142 expander (turbine). Process 1-2 is isosteric heating same as in BCAS, while process 2-3 is expansion  
143 process from the bed pressure (maximum pressure) to the condenser pressure, process 3-4 is isobaric  
144 condensation while the rest processes are similar to BCAS.

145



146  
147

Figure 1: Schematic diagram of (a) Basic cooling adsorption system (BCAS) (b) Adsorption system for cooling and electricity.



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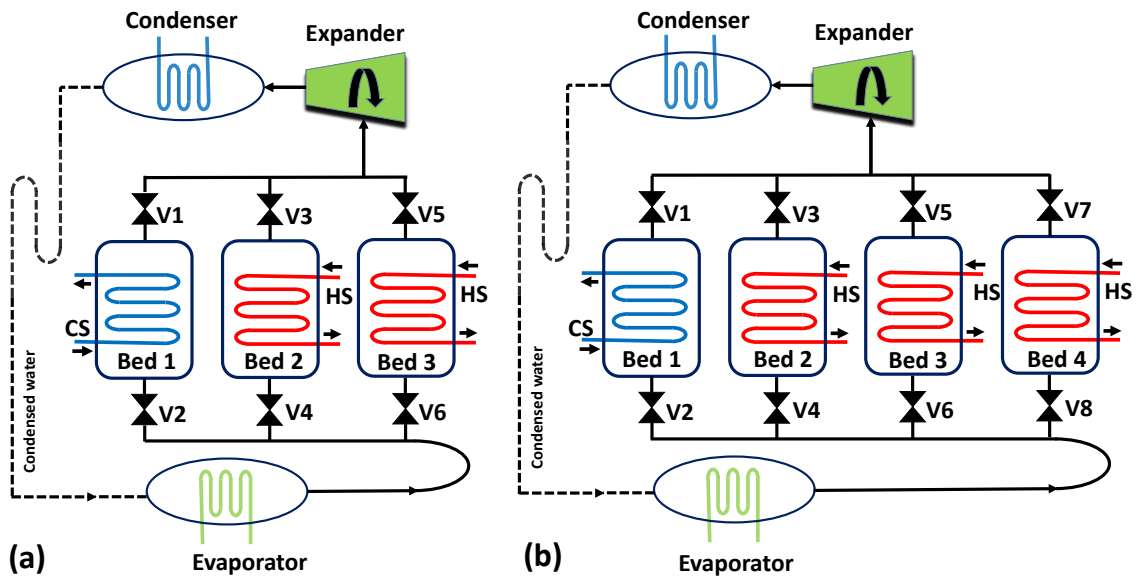
Figure 2: P-T diagram (a) Basic cooling adsorption system (BCAS) (b) Adsorption system for cooling and electricity.

151 **3. Multi-bed adsorption system for cooling and electricity**

152 Multi-bed adsorption system for cooling and electricity has the same principle of work for the two-bed  
 153 adsorption system for cooling and electricity as discussed in section 2. Instead of using only two  
 154 adsorption beds, a number of adsorption beds are used in the same system. Usually when two-bed is used  
 155 in the adsorption system the total adsorption time (adsorption time +switching time) is equal to the total

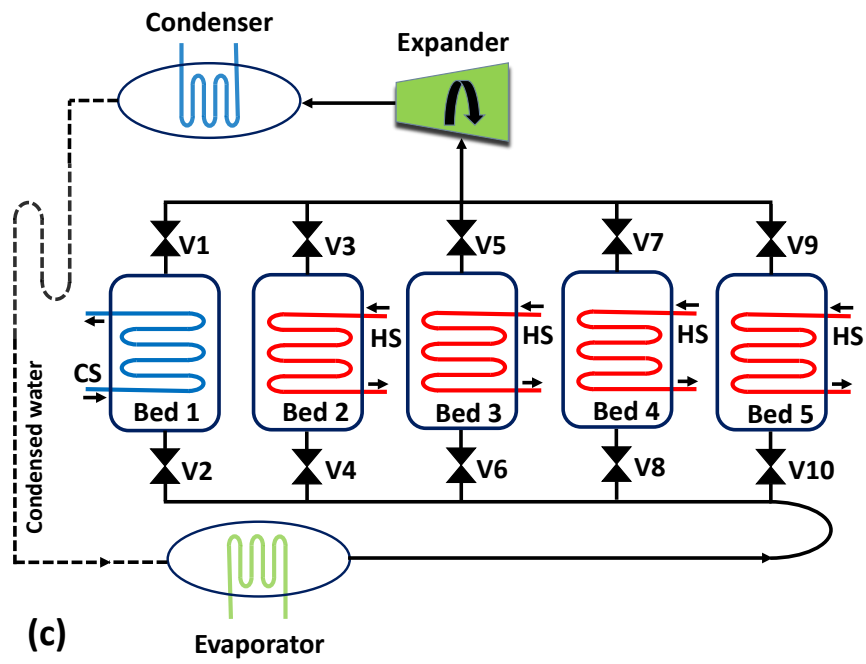
156 desorption time (desorption time + switching time) .i.e. the time ratio  $R=1$ . This study aims to investigate  
 157 the effect of using multi-bed configurations with a number of time ratios ( $R$ ) on the overall system  
 158 performance. For example, in the three-bed adsorption system with  $R=1/2$  shown in Figure 3 (a), bed 1  
 159 starts with desorption phase (20s switching time + 140s net desorption) as shown in Figure 4 with zero  
 160 delay time, then it switches to adsorption phase (20s switching time + 300s net adsorption). Bed 2 and  
 161 bed 3 work in similar way as shown in Figure 4, but with delay times of 160s and 320s respectively. The  
 162 same concept can be used for the four, five and six-beds systems as shown in Figure 3 (b), (c) and (d)  
 163 with  $R$  of  $1/3$ ,  $1/4$  and  $1/5$  respectively.

164 Besides two (2B), three (3B), four (4B), five (5B) and six-bed (6B) systems another two configurations  
 165 are investigated in this study. The first one is the four-bed system which consisting of a pair of two beds  
 166 working in parallel with  $R=1$  as shown in Figure 3(e), while the other one is six-bed which consisting of a  
 167 pair of three beds working in parallel with  $R=1/2$  as shown in Figure 3(f). Thus in this work, 9 different  
 168 cases including 7 different bed configurations and 7 adsorption/desorption ratios are investigated to find  
 169 the best configuration and adsorption/desorption ratio in terms of performance, specific cooling and  
 170 specific power output.

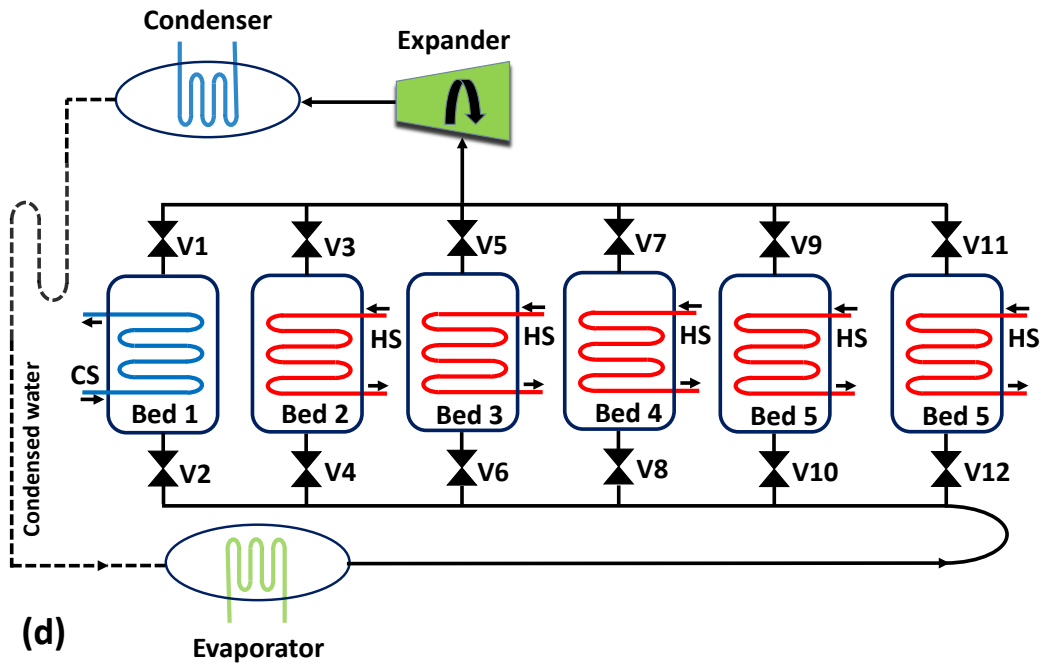


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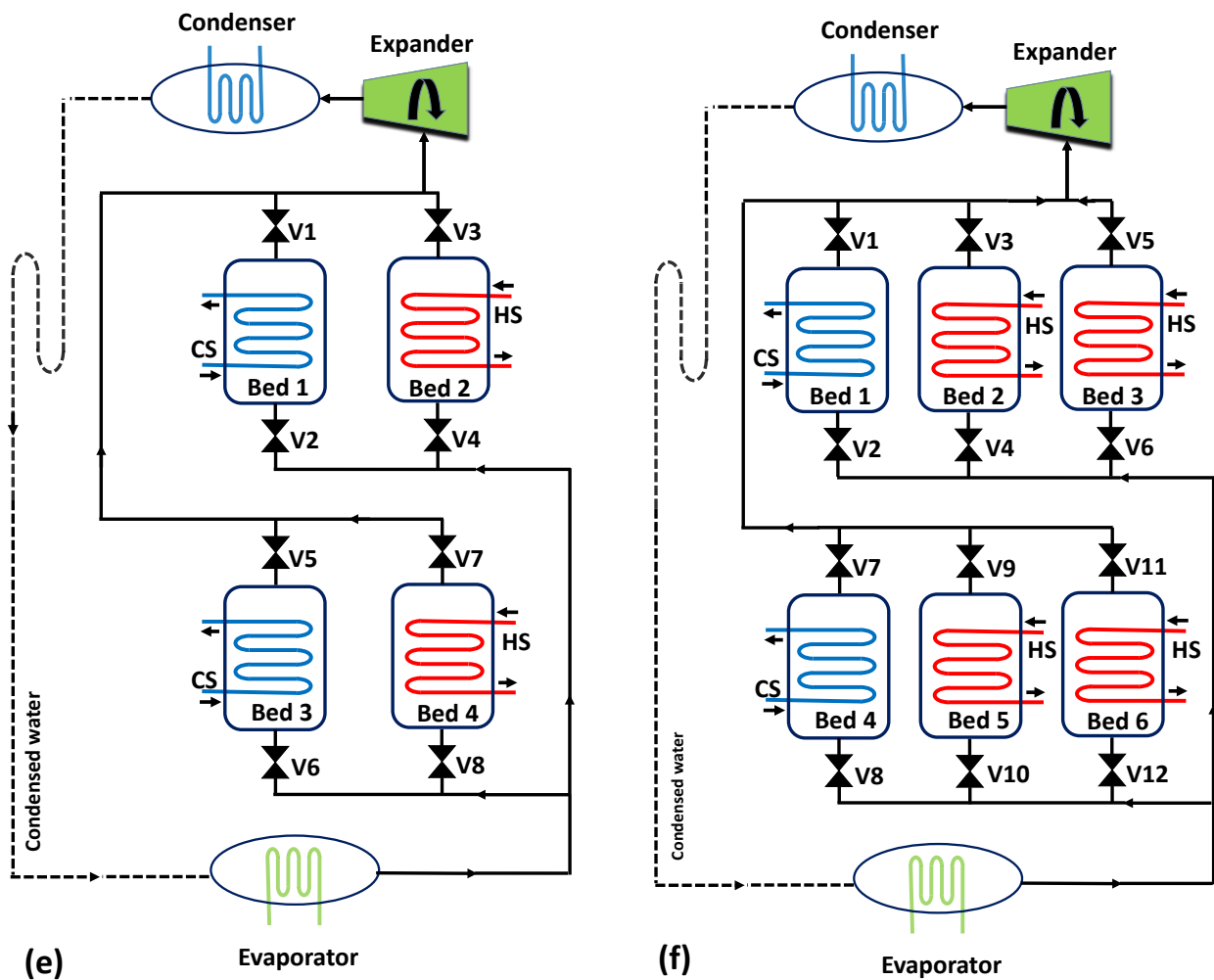


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175  
 176 Figure 3: Schematic diagram of multi bed adsorption system for cooling and electricity (a) three-bed (b) four-bed in series  
 177 (c) five-bed (d) six-bed in series (e) four-bed in parallel and (f) six-bed in parallel.

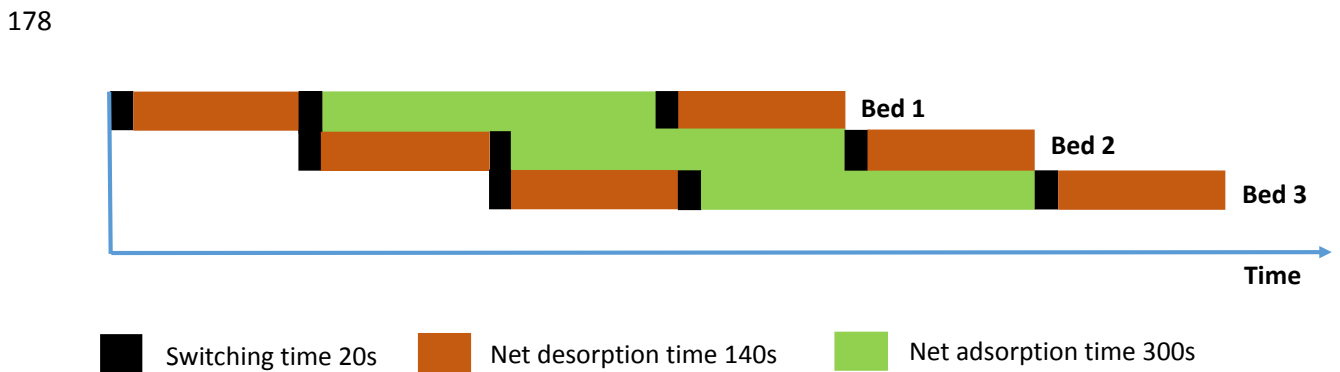


Figure 4: Time allocation of the three-bed adsorption system for cooling and electricity with  $R=1/2$ .

185 **4. Mathematical Modelling**

186 In this study, new advanced adsorption pairs of AQSOA-ZO2/water and Aluminium-Fumarate/water are  
 187 used and compared to Silica-gel/water. The adsorption equilibrium for AQSOA-ZO2/water can be written  
 188 as [37]

189 
$$x_{eq} = x_o \left[ \frac{k \left( \frac{P_s(T_w)}{P_s(T_a)} \right)^n}{1 + (k-1) \left( \frac{P_s(T_w)}{P_s(T_a)} \right)^n} \right] \quad (1)$$

190 
$$k = \alpha \exp \left[ n(Q_{st} - h_{fg}) / (RT^{bed}) \right] \quad (2)$$

191 Where,  $P_s(T_w)$  and  $P_s(T_a)$  are the saturation vapour pressure at water vapour temperature and adsorbent  
 192 temperature respectively and the constants  $x_o$ ,  $\alpha$ ,  $n$  and  $Q_{st}$  are taken from [26].

193 For Aluminium-Fumarate/water, the adsorption equilibrium of equation (3) is obtained from [38] and  
 194 listed in Table 1.

195 
$$x_{eq} = f(A) \quad (3)$$

196  
 197 *Where*

198 
$$A = -RT^{bed} \ln \left( \frac{P_s(T_w)}{P_s(T_a)} \right) \quad (4)$$

199 **Table 1: Aluminium-Fumarate/water isotherms of Eq. (3)**

$A < 2900$	$x_{eq} = 0.5948 - 3.12E-4A + 1.68302E-7A^2 - 3.124455E-11A^3$
$A > 3987$	$x_{eq} = 0.111993 \text{EXP}(-0.000258797A)$
$2900 \leq A \leq 3987$	$x_{eq} = 2.36129 - 9.93768E-4A + 1.05709E-7A^2$

200  
 201 For Silica-gel/water, the modified Freundlich equation is used to present the adsorption equilibrium  
 202 [39-41] as:

203 
$$x_{eq} = A(T_s) \left[ \frac{P_s(T_w)}{P_s(T_a)} \right]^{B(T_s)} \quad (5)$$

204 *where*

205 
$$A(T_s) = A_o + A_1 T_s + A_2 T_s^2 + A_3 T_s^3 \quad (6)$$

206 
$$B(T_s) = B_o + B_1 T_s + B_2 T_s^2 + B_3 T_s^3 \quad (7)$$

207 The constants of equations (6) and (7) are obtained from [41, 42].

208 Linear driving force (LDF) equation is used to describe the adsorption/desorption rate as [39, 40, 43]

209 
$$\frac{dx}{dt} = k_o \exp \left( -\frac{E_a}{RT} \right) (x_{eq} - x) \quad (8)$$

210 The kinetics constants ( $k_o$  and  $E_a$ ) of equation (8) are taken from reference [44] for AQSOA-ZO2/water,  
 211 from reference [38] for Aluminium-Fumarate/water and from references [39, 40] for Silica-gel/water.

212 Lumped model method is used define the energy balance in adsorption beds, where the adsorbent, the  
 213 adsorbate and the bed materials are assumed to be at the same temperature during the cycle time [43, 45,  
 214 46].

$$215 \quad \left( M c_{p, \text{eff}}^{\text{bed}} \right) \frac{dT^{\text{bed}}}{dt} + \left( M_a x_i^{\text{bed}} c_p \right) \frac{dx_i^{\text{bed}}}{dt} = \varphi M_a \left( \frac{dx_i^{\text{bed}}}{dt} \right) (Q_{st}) - (\dot{m} c_p)_j (T_{j,o} - T_{j,in}) \quad (9)$$

216 Flag  $\varphi$  equals to 0 at switching time and equals to 1 at adsorption/desorption process and the bed outlet  
 217 temperature is given by [43, 45].

$$218 \quad T_{j,o} = T_i^{\text{bed}} + (T_{j,in} - T_i^{\text{bed}}) \exp \left[ \frac{-(U A_r)_i^{\text{bed}}}{(\dot{m} c_p)_j} \right] \quad (10)$$

219 The energy balance equations for the condenser can be expressed by [46, 47].

$$220 \quad \left( M c_{p, \text{eff}}^{\text{cond}} \right) \frac{dT^{\text{cond}}}{dt} = \varphi h_{fg} M_a \frac{dx_{\text{des}}^{\text{bed}}}{dt} - (\dot{m} c_p)_{\text{cond}} (T_{w,o} - T_{w,i}) - (c_p)_w (T^{\text{bed}} - T^{\text{cond}}) M_a \frac{dx_{\text{des}}^{\text{bed}}}{dt} \quad (11)$$

221 The condenser outlet temperature is given by [46, 47].

$$222 \quad T_{w,o} = T^{\text{cond}} + (T_{w,in} - T^{\text{cond}}) \exp \left[ \frac{-(U A_r)^{\text{cond}}}{(\dot{m} c_p)_{\text{cond}}} \right] \quad (12)$$

223 The energy balance in the evaporator is expressed as [46, 47].

$$224 \quad \left( M c_{p, \text{eff}}^{\text{evap}} \right) \frac{dT^{\text{evap}}}{dt} = \varphi h_{fg} M_a \frac{dx_{\text{ads}}^{\text{bed}}}{dt} - (\dot{m} c_p)_{\text{evap}} (T_{\text{chill},o} - T_{\text{chill},i}) - (c_p)_w (T^{\text{cond}} - T^{\text{evap}}) M_a \frac{dx_{\text{des}}^{\text{bed}}}{dt} \quad (13)$$

225 The outlet temperature of the chilled water can be written as [40, 46, 47].

$$226 \quad T_{\text{chill},o} = T^{\text{evap}} + (T_{\text{chill},in} - T^{\text{evap}}) \exp \left[ \frac{-(U A_r)^{\text{evap}}}{(\dot{m} c_p)_{\text{evap}}} \right] \quad (14)$$

227 The mass balance in the adsorption evaporator can be written as [40, 43, 45, 46].

$$228 \quad \frac{dM_{\text{ref}}}{dt} = -M_a \left[ \frac{dx_{\text{des}}^{\text{bed}}}{dt} + \frac{dx_{\text{ads}}^{\text{bed}}}{dt} \right] \quad (15)$$

229 Mechanical work produced in the expander (turbine), can be written as below:

$$W_{exp} = \frac{\int_0^{t_{cycle}} \eta_{exp} \dot{m}_{ads} \Delta h dt}{t_{cycle}} \quad (16)$$

Where  $\Delta h$  is the enthalpy difference through the adsorption expander and  $\dot{m}_{ads}$  is the water mass flow rate through the expander, while  $\eta_{exp}$  is the expansion efficiency which is assumed to be ideal for this thermodynamic study. The overall performance of the adsorption system for cooling and electricity can be defined using the terms SCP, SP, COP, adsorption power efficiency ( $\eta_{ads}$ ), COPe and SCPe and written in equations (17-21).

$$SCP = \frac{(\dot{m}_{cp})_{evap} \int_0^{t_{cycle}} (T_{chill,o} - T_{chill,i}) dt}{M_a t_{cycle}} \quad (17)$$

$$SP = \frac{\int_0^{t_{cycle}} \eta_{exp} \dot{m}_{ads} \Delta h dt}{M_a t_{cycle}} \quad (18)$$

$$\eta_{ads} = \frac{\int_0^{t_{cycle}} \eta_{exp} \dot{m}_{ads} \Delta h dt}{(\dot{m}_{cp})_h \int_0^{t_{cycle}} (T_{h,o} - T_{h,i}) dt} \quad (19)$$

$$COPe = \frac{(\dot{m}_{cp})_{evap} \int_0^{t_{cycle}} (T_{chill,o} - T_{chill,i}) dt + F [\int_0^{t_{cycle}} \eta_{exp} \dot{m}_{ads} \Delta h dt]}{(\dot{m}_{cp})_h \int_0^{t_{cycle}} (T_{h,o} - T_{h,i}) dt} \quad (20)$$

240

$$SCPe = \frac{(\dot{m}_{cp})_{evap} \int_0^{t_{cycle}} (T_{chill,o} - T_{chill,i}) dt + F [\int_0^{t_{cycle}} \dot{m}_{ads} \Delta h dt]}{M_a t_{cycle}} \quad (21)$$

The term ( $\eta_{ads}$ ) is used to represent the power generation efficiency of the adsorption system i.e. the ratio of the amount of power generated through the expander (turbine that incorporated within the adsorption system) to the total heat consumed by the adsorption beds. The terms *COPe* (equivalent coefficient of performance) and *SCPe* (equivalent specific cooling power) are used to compare the performance of adsorption system for cooling and electricity to the two-bed basic cooling adsorption system (BCAS). *F* is the typical COP for compression refrigeration system, which is assumed to be (3) in this work i.e. the power generated, by adsorption system can be converted into cooling again and in order to compare between adsorption system for cooling and electricity system and the basic cooling adsorption system (BCAS).

251 The Exergy efficiency depending on the second law of thermodynamic can be defined as the ratio  
 252 between the exergy output to the exergy input and it is used to highlight the different grade of cooling and  
 253 electricity generated by the adsorption system used in this paper. The exergy efficiency can be defined as  
 254 [17, 48]:

$$255 \quad \eta_{ex} = \frac{W_{exp} + E_{evap}}{E_{in}} \quad (22)$$

256 Where  $E_{evap}$  is the cooling exergy through the evaporator and can be defined as [49-51]:

$$257 \quad E_{evap} = \frac{(\dot{m}c_p)_{evap} \int_0^{t_{cycle}} (T_{chill,o} - T_{chill,i}) dt}{t_{cycle}} \left[ \frac{T^{amb}}{T_{evap}} - 1 \right] \quad (23)$$

258 While,  $E_{in}$  is the exergy input to the system and can be defined as [17, 48]:

$$259 \quad E_{in} = \dot{m}_h [(h_{h,in} - h_{h,o}) - T^{amb} (s_{h,in} - s_{h,o})] \quad (24)$$

## 260 5. Results and discussions

261 Table 2 (a) shows the main operating conditions used in this work, while Table 2 (b) and (c) show the  
 262 features of the main components characteristics (bed, condenser, and evaporator) used in this study.  
 263 Figure 5 shows the values of COP/COPE, SCP/SCPE and exergy efficiency for two-bed basic cooling  
 264 adsorption system (BCAS) and adsorption system for cooling and electricity (with two-bed configuration)  
 265 for Silica-gel, AQSOA-Z02, and Aluminium-Fumarate with a range of heat source temperature. For most  
 266 cases, COPE achieved by the adsorption system for cooling and electricity using Silica-gel, Aluminium-  
 267 Fumarate, and AQSOA-Z02 is higher than the COP of BCAS and this is because additional electricity is  
 268 generated in adsorption system for cooling and electricity. However, for AQSOA-Z02 with heat source  
 269 temperature of 120 °C (or less), the COP of BCAS is higher than COPE of adsorption system for cooling  
 270 and electricity and this is due to AQSOA- Z02 shows low performance with adsorption system for  
 271 cooling and electricity at low heat source temperatures. Moreover, SCPE achieved by the adsorption  
 272 system for cooling and electricity is higher than that produced by BCAS for most cases using Silica-gel,  
 273 Aluminium-Fumarate, and AQSOA-Z02 (except for Silica-gel at 80 °C and AQSOA-Z02 at 120 °C or  
 274 less and this is because less power is generated with low heat source temperatures). At high heat source  
 275 temperatures the mass flow rate of refrigerant (water) is higher because of high adsorption/desorption

276 rate. In addition, at high heat source temperatures, high pressure ratio can be obtained through the  
277 expander (turbine) which means more electricity can be generated. As the grade of electricity is higher  
278 than cooling (i.e. each 1kW of electricity produces about 3 kW of cooling depending on the typical COP  
279 for compression refrigeration system, which is assumed to be three in this study), SCPe of adsorption  
280 system for cooling and electricity is higher than that of BCAS at heat source temperature higher than 120  
281 °C. For all adsorption materials and heat source temperatures used, the exergy efficiency achieved by  
282 adsorption system for cooling and electricity is higher than that for BCAS and this is because the electric  
283 power generated by the former has high grade than cooling. The maximum exergy efficiency of 54% is  
284 achieved using Silica-gel at heat source temperature of 80 °C, results also show that using adsorption  
285 system for cooling and electricity can enhance the exergy efficiency of BCAS by up to 2.5 times when  
286 using Al-Fumarate at heat source temperature of 160 °C. Also, results showed that, different adsorption  
287 materials presented different values of COP/COPE, SCP/SCPe, and exergy efficiency. For example,  
288 Silica-gel showed the highest COP and exergy efficiency, while AQSOA-Z02 showed the highest SCP  
289 with heat source temperature of 140 °C or higher. COP is the most important coefficient of any heat  
290 pump, because high COP values means less energy used. However, if the energy used is infinite or semi-  
291 infinite source like solar energy or geothermal energy, SCP can be the most important criterion, because  
292 high SCP means more cooling is generated using the same heat pump size. For system generating cooling  
293 and electricity, exergy efficiency is essential because the electricity has different grade compared to  
294 cooling, so it helps to make a good comparison between BCAS system and the adsorption system for  
295 cooling and electricity.

296 Figure 6, Figure 7, and Figure 8 show the cooling and power (electricity) generated using 7 different  
297 configurations utilizing Silica-gel, AQSOA-Z02 and Aluminium-Fumarate with heat source temperature  
298 of 120°C. Results show that cooling and electricity can be generated at the same time however, the  
299 amount of cooling and electricity generated is varied from one configuration to another and from one  
300 material to another. As the number of beds increases the amount of cooling and electricity increases  
301 because of more adsorption materials are added to the system and more uptake and mass flow can be  
302 generated. In addition, as the number of beds increases, more continuity in cooling and electricity can be  
303 noticed i.e. the cooling and electricity with less fluctuation which is more preferable. A significant gain in  
304 cooling and electricity can be noticed between 2B (two-beds) and 3B (three-bed) configurations where

305 more cooling and electricity can be generated with configuration 3B. Configuration 6B (six-beds) in  
306 parallel with  $R=1/2$  gives the highest average cooling and electricity generated as shown Figure 6, Figure  
307 7, and Figure 8 for Silica-gel, AQSOA-Z02, and Aluminium-Fumarate respectively.

308 Figure 9 shows the COP of different configurations and R ratios utilizing Silica-gel, AQSOA-Z02, and  
309 Aluminium-Fumarate with a range of heat source temperature between 80 and 160 °C. The maximum  
310 COP achieved in this investigation is 0.64 using Silica-gel at heat source temperature of 80 °C with 2B  
311 configuration and  $R=1$ . Compared to other materials at heat source temperature between 80-120 °C,  
312 Silica-gel shows the highest COP and this is due to the high cooling capacity achieved with this material  
313 as a result of high water uptake (high adsorption/desorption rate). Silica-gel's isotherms which has linear  
314 and uniform shape, besides its good kinetics helps to generate such high uptake rate and this can explain  
315 the high water and the high cooling capacity produced by this material. Regarding the heat source  
316 temperature, at 80 °C the amount of heat consumed is the lowest which leads to highest COP. In terms of  
317 the number of beds and R ratio used, the configuration 2B with  $R=1$  shows the maximum COP and this is  
318 due to less amount of heat when using 2B configuration.

319 Figure 10 shows SCP of different configurations and R ratios utilizing Silica-gel, AQSOA-Z02 and  
320 Aluminium-Fumarate with heat source temperature between 80 and 160 °C. The maximum SCP achieved  
321 is around 650 W/kg<sub>ads</sub>. Using AQSOA-Z02 at heat source temperature of 160 °C with three-bed  
322 configuration and  $R=1/2$ . Generally, desorption rate is faster than adsorption rate [32] and this is because  
323 the later occurs at relatively low temperature (28 °C in this study as an example), so discharging the  
324 refrigerant (water) from the adsorption material can be faster than charging the materials with the  
325 refrigerant. As a result, there is an optimum R ratio for each case depending on the adsorption material  
326 and the regeneration temperature used. Again, this can explain why the configurations with  $R>1$  have low  
327 SCP compared to other configurations, as the time of adsorption is more than that of desorption.

328 Figure 11 shows SP of different configurations and R ratios utilizing Silica-gel, AQSOA-Z02 and  
329 Aluminium-Fumarate with heat source temperature between 80 and 160 °C. Maximum SP generated in  
330 this study is about 64 W/kg utilizing AQSOA-Z02 at 160 °C with 3B configuration and  $R=1/2$ . SP of 64  
331 W/kg<sub>ads</sub> is not very large value compared to the value of cooling (650 W/kg<sub>ads</sub>), however the grade of  
332 electricity is higher than that of cooling because ideally each 1kW of electricity can generate 3kW of  
333 cooling when the typical COP for compression refrigeration system is assumed to be 3. In terms of the



334 number of beds and R ratio used, the configuration 3B with  $R=1/2$  has the maximum SP for Silica-gel and  
335 Aluminium-Fumarate for all the range of heat source temperatures used, while for AQSOA-Z02 this  
336 occurs only with heat source temperature of 160 °C and this can show that there is a specific limit of heat  
337 source temperature, where after such limit the 3B configuration can be the best. The configurations with  
338  $R>1$  have low SP compared to other configurations, as the time of adsorption is more than that of  
339 desorption.

340 Figure 12 shows adsorption power efficiency of different configurations and R ratios utilizing Silica-gel,  
341 AQSOA-Z02, and Aluminium-Fumarate with heat source temperature between 80 and 160 °C. The  
342 maximum power efficiency achieved in this study is 4.63 % using AQSOA-Z02 at heat source  
343 temperature of 160 °C with 2B configuration and  $R=1$ . As the heat source temperature increases, the mass  
344 flow rate through the expander (turbine) increases because of high desorption rate. Moreover, high heat  
345 source temperature produces more power and then high adsorption power efficiency and this is mainly  
346 because the high pressure difference between the desorber and the condenser.

347 The main advantage of using multi-bed adsorption system for cooling and electricity is to increase the  
348 values of SCP and SP generated from this system and offer better options to users and designers. Besides  
349 increasing the quantity of the cooling and electricity, they can be generated with less fluctuation and more  
350 continuity. However, using multi-bed adsorption configurations may lead to large size and heavy weight,  
351 but this problem can be solved by developing new adsorption materials with better adsorption capacity,  
352 and improving the design of the bed heat exchanger, besides using lighter and more efficient materials  
353 in the bed heat exchanger. Also, this study shows that four-bed configuration consisting of a pair of two-  
354 beds working in parallel produces more COP and SCP than the configuration with a same number of beds  
355 working in series. The same result can be noticed for the six-bed configuration working in parallel  
356 compared to the same number of beds working in series and this is due to R ratio used in each  
357 configuration of  $R=1/2$  and  $R=1/3$  which produce the highest COP and SCP respectively.

358

**Table 2:**

359

**a) Parameters used in simulation**

<b>Parameter</b>	<b>Value</b>
Ambient temperature °C	36
Bed heating fluid temperature °C	80-140
Bed cooling fluid temperature °C	23-61
Condenser cooling temperature °C	28
Chilled water temperature °C	18
Bed hot fluid mass flow rate kg/s	1.7
Bed cold fluid mass flow rate kg/s	1.6
Condenser mass flow rate kg/s	0.363
Evaporator mass flow rate kg/s	0.8
Adsorption /desorption phase times (2 bed)	300+20
Switching time s	28-64

366

**b) Bed heat exchanger characteristics [26]**

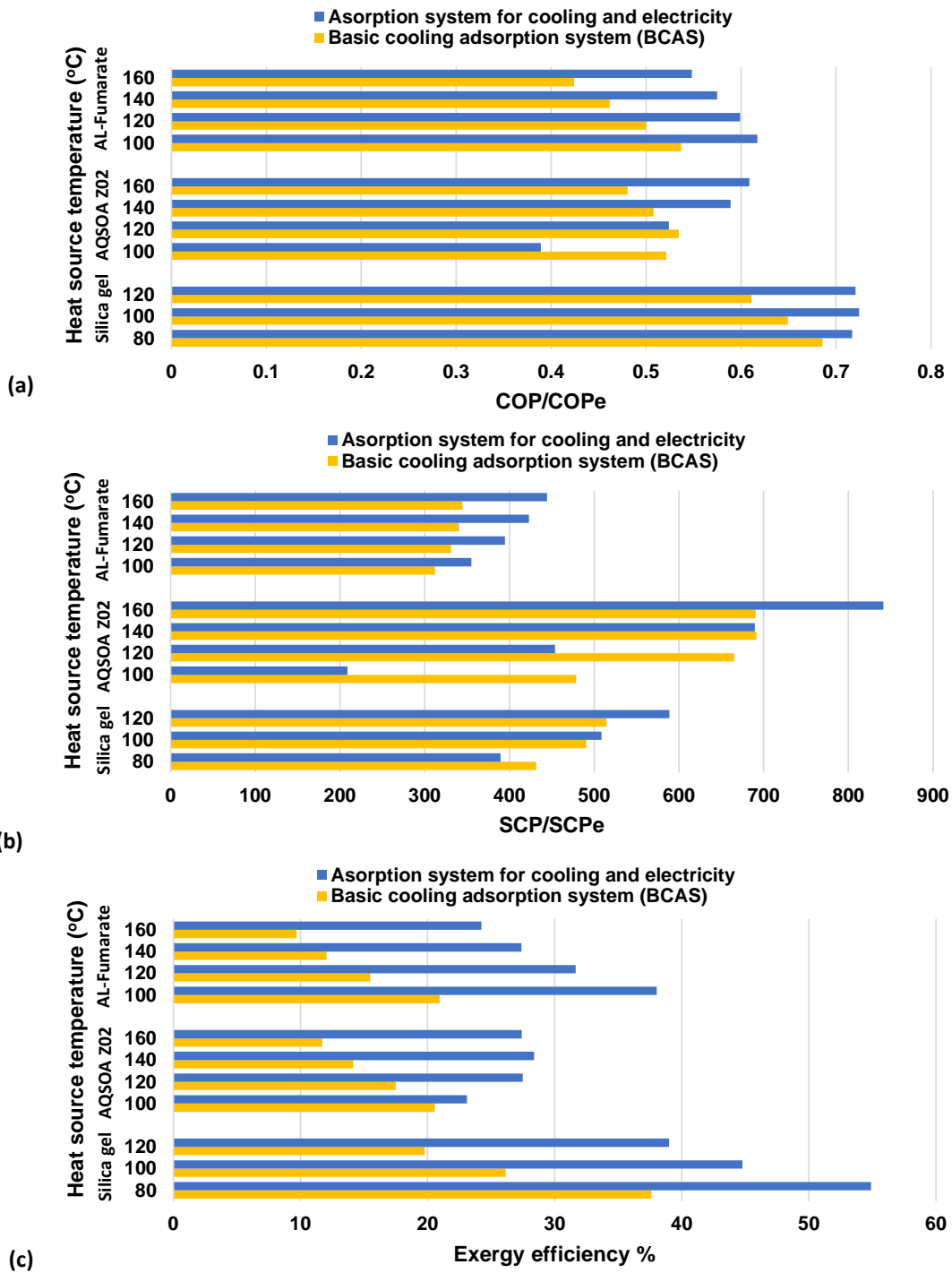
<b>Parameter</b>	<b>Value</b>
Fin length m	172E-3
Fin width m	30E-3
Fin pitch m	1.2E-3
Module length m	450E-3
Finned length m	370E-3
No. of module	4
No. tubes/module	6
Tube OD m	15.875E-3
Tube thickness m	0.8E-3

367

**c) Condenser/evaporator characteristics [26]**

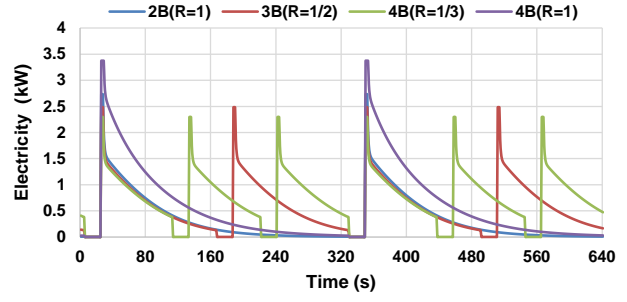
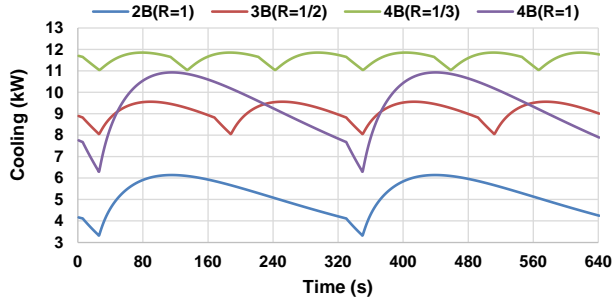
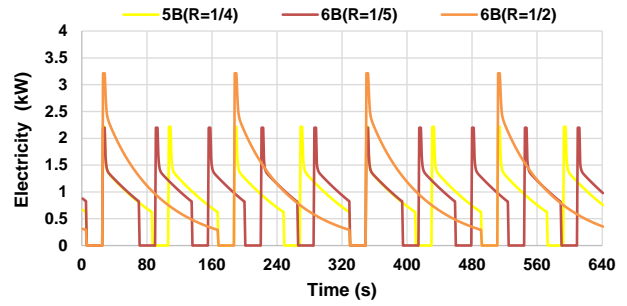
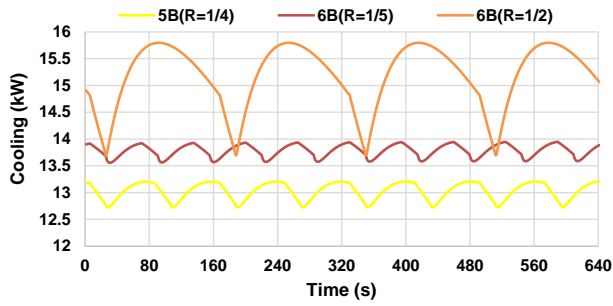
<b>Parameter</b>	<b>Value</b>
Pipe length m	5.5
No. tubes	4
Tube OD m	15.875E-3
Tube thickness m	0.8E-3

368



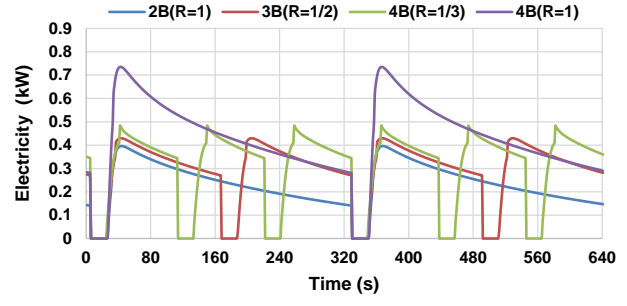
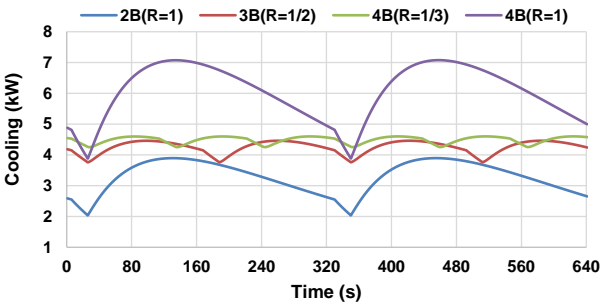
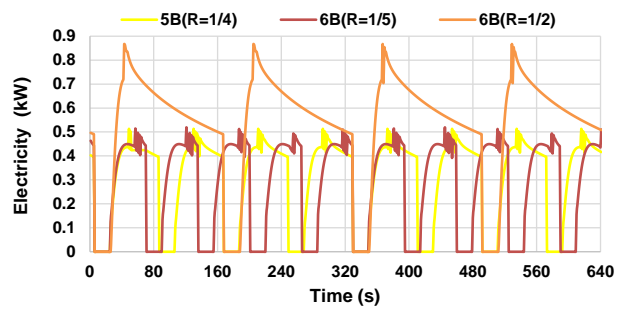
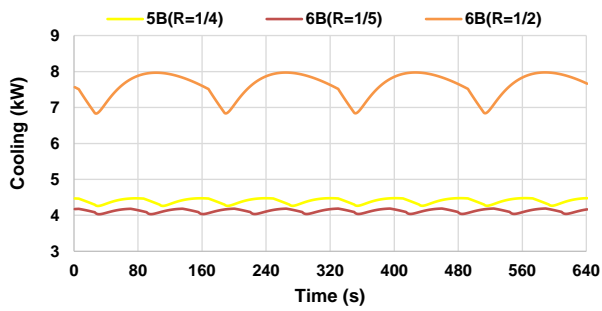
369  
 370 **Figure 5: COP/COPe, SCP/SCPe and Exergy efficiency of basic cooling adsorption system (BCAS) and adsorption system**  
 371 **for cooling and electricity (with two bed) for a range of heat source temperature utilizing Silica-gel, AQSOA-Z02 and**  
 372 **Aluminium-Fumarate.**

373



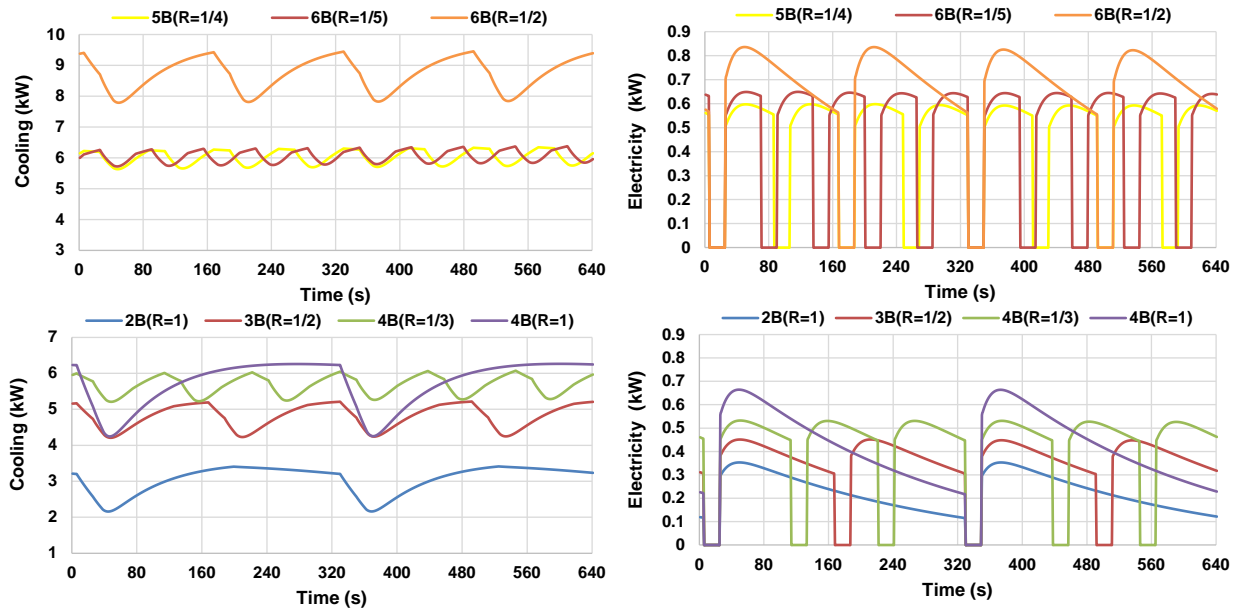
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Figure 6: Cooling and electricity generated from multi-bed adsorption system utilizing Silica-gel at heat source temperature of 120 °C.



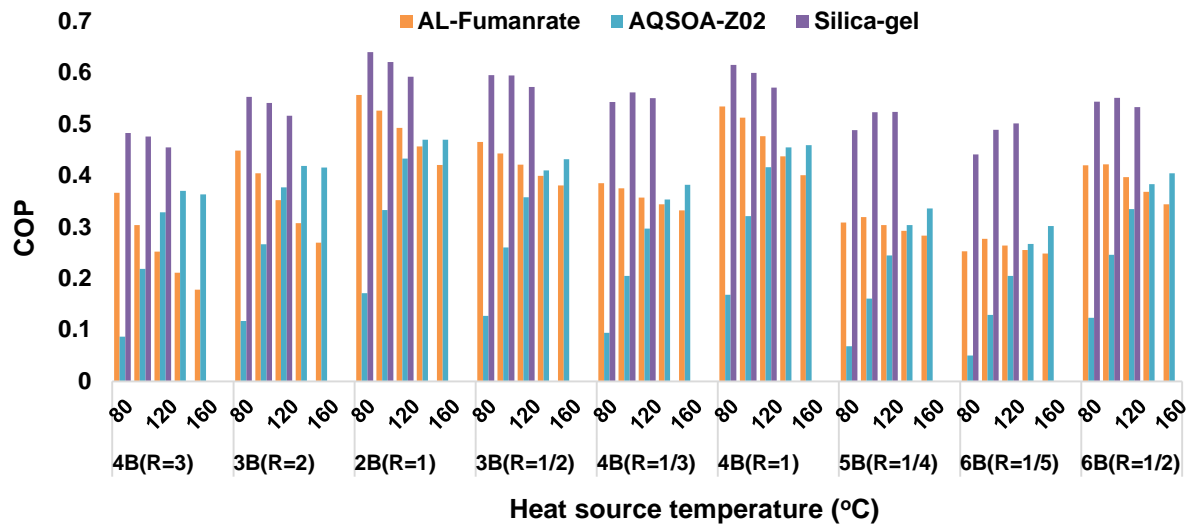
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Figure 7: Cooling and electricity generated from multi-bed adsorption system utilizing AQSOA-Z02 at heat source temperature of 120 °C.



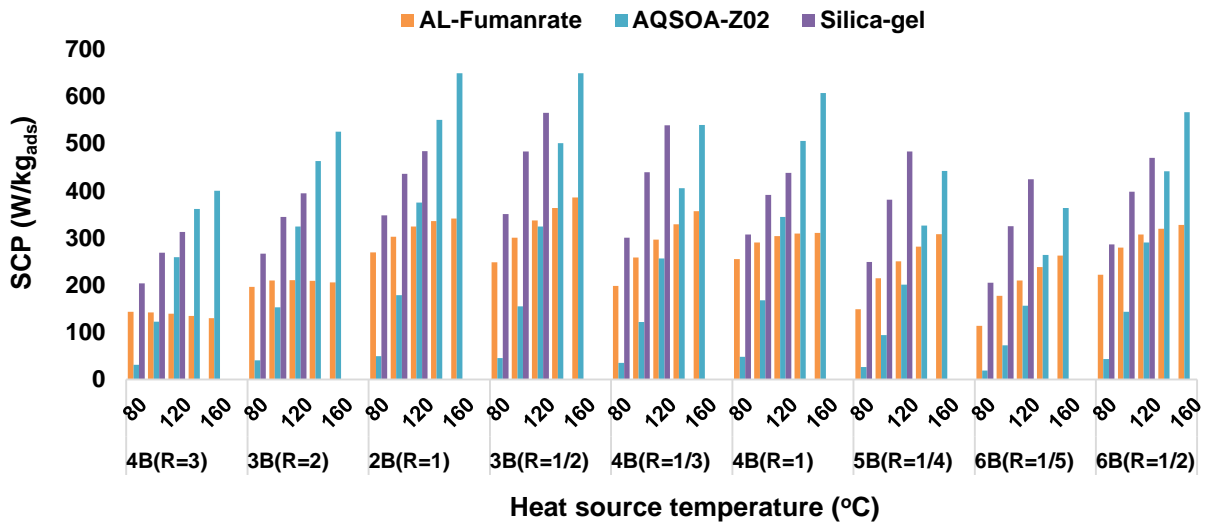
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Figure 8: Cooling and electricity generated from multi-bed adsorption system utilizing Aluminium-Fumarate at heat source temperature of 120 °C.

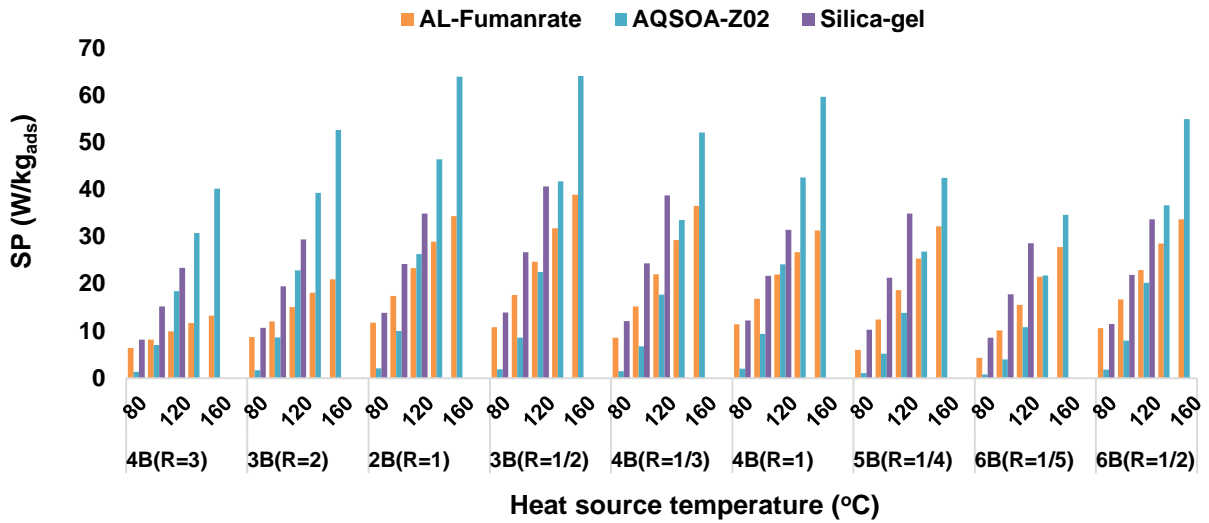


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Figure 9: COP of different configurations of multi-bed adsorption system and adsorption/desorption ratio (R) utilizing Silica-gel, AQSOA-Z02 and Aluminium-Fumarate with a range of heat source temperature.

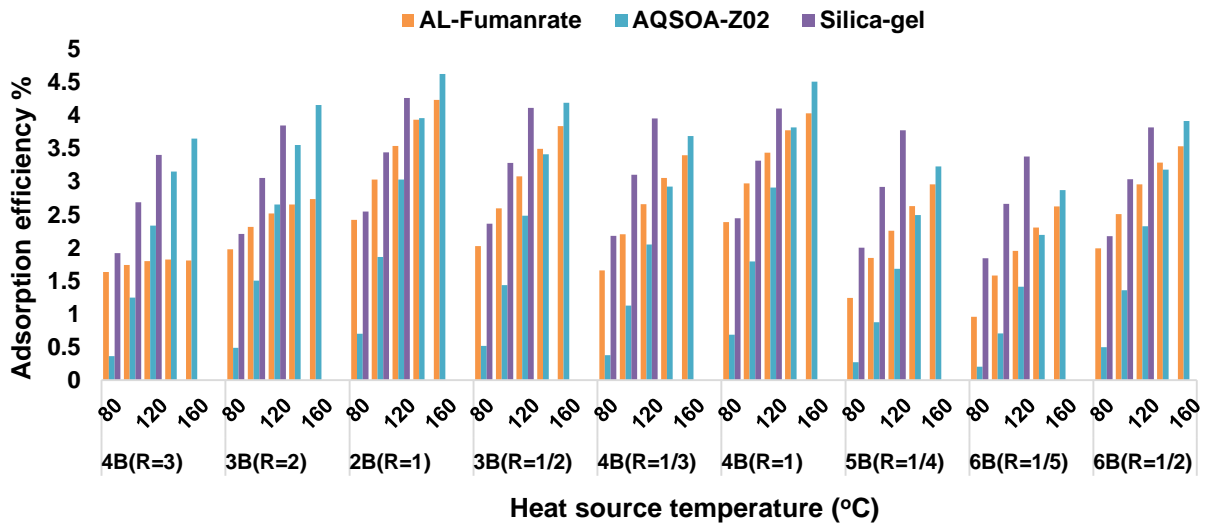


388  
 389 **Figure 10: SCP of different configurations of multi-bed adsorption system and adsorption/desorption ratio (R) utilizing**  
 390 **Silica-gel, AQSOA-Z02 and Aluminium-Fumarate with a range of heat source temperature.**  
 391



392  
 393 **Figure 11: SP of different configurations of multi-bed adsorption system and adsorption/desorption ratio (R) utilizing**  
 394 **Silica-gel, AQSOA-Z02 and Aluminium-Fumarate with a range of heat source temperature.**  
 395

396



398 **Figure 12: Adsorption power efficiency of different configurations of multi-bed adsorption system and**  
399 **adsorption/desorption ratio (R) utilizing Silica-gel, AQSOA-Z02 and Aluminium-Fumarate with a range of heat source**  
400 **temperature.**

## 401 6. Conclusions

402 In this paper, 9 cases including 7 multi-bed adsorption system configurations and 7 different  
403 adsorption/desorption time ratios for generating cooling and electricity at the same time have been  
404 designed and simulated. Furthermore, advanced adsorption materials of AQSOA-Z02/water,  
405 Aluminium-Fumarate MOF/water, and Silica-gel/water are studied in terms of system performance  
406 and compared to each other. The main results of this study can be listed as:

- 407
- 408 1. Adsorption system for cooling and electricity is feasible and can generate cooling and electricity  
409 simultaneously. For example, utilizing 10.76 kg of Silica-gel at 120 °C of heat source temperature  
410 can generate an average cooling of 15.18 kW and an average electricity of 0.94 kW.
- 411 2. Adsorption system for cooling and electricity has higher COPE than basic cooling adsorption system  
412 (BCAS) except for AQSOA-Z02 with heat source temperature below 120 °C.
- 413 3. For all adsorption material and heat source temperatures used in this study, the exergy efficiency of  
414 adsorption system for cooling and electricity is higher than that for BCAS, and maximum exergy  
415 efficiency of 54% is achieved utilising Silica-gel at 80 °C.
- 416 4. Adsorption system for cooling and electricity has higher SCPE than BCAS for Silica-gel with heat  
417 source temperature higher than 100 °C, for AQSOA-Z02 with heat source temperature higher than  
418 140 °C and for Aluminium-Fumarate with all heat source temperature used in this study.
- 419 5. Two-bed configuration with R=1 (adsorption/desorption time ratio =1) has the maximum COP for  
420 all adsorption materials and all range of temperatures used in this work.

- 421 6. Three-bed configuration with  $R=1/2$  has the maximum SCP and SP for Silica-gel and Aluminium-  
 422 Fumarate for all the range of heat source temperatures used, while for AQSOA-Z02 with heat  
 423 source temperature of 160 °C.
- 424 7. As the number of beds increases more than two, the COP decreases, while as the number of beds  
 425 increases more than two or three (depending on materials and heat source temperature used) the SCP  
 426 and SP decrease.
- 427 8. As the number of bed increases, more continuity in cooling and electricity can be achieved.
- 428 9. Pairs of two-bed (four beds) and pairs of three-bed (six beds) configurations working in parallel  
 429 produce more COP and SCP than four-bed and six-bed configurations working in series.

430 **Nomenclature**

<b>Symbols</b>			
$A$	adsorption potential, J/mole	$\eta$	efficiency
$Al$	aluminium	$\alpha$	constant used in eq. 2
$A_r$	area, $m^2$	$\rho$	density $kg/m^3$
$C_p$	specific heat capacity, J/kg.K	$\varphi$	flag
<b>COP</b>	coefficient of performance	<b>Subscript</b>	
<b>COPe</b>	equivalent coefficient of performance	<i>ads,a</i>	adsorbent
$E$	Exergy kW	<i>ads</i>	adsorption
$k_o$	empirical constant in Eq. (6), 1/s	<i>amb</i>	ambient temperature
$E_a$	activation energy, J/kg	<i>bed</i>	adsorbent bed
$h$	enthalpy, J/kg	<i>chill</i>	chilled water
$h_{fg}$	evaporation latent heat J/kg	<i>cond</i>	condenser
$M$	mass, kg	<i>des</i>	desorption
$\dot{m}$	mass flow rate, kg/s	<i>eff</i>	effective
$P$	pressure, Pa	<i>evap,e</i>	evaporator
$Q_{st}$	isosteric heat of adsorption, J/kg	<i>exp</i>	expander
$R$	gas constant (for water vapour), J/kg.K	<i>ex</i>	exergy
$R$	adsorption/desorption time ratio	<i>f</i>	liquid
$U$	overall heat transfer coeff., $W/m^2K$	<i>g</i>	gas
$W$	power generated W	<i>i</i>	adsorption/desorption
$SP$	specific power generated $W/kg_{ads}$	<i>in</i>	inlet
$SCP$	specific cooling power $W/kg_{ads}$	<i>j</i>	cooling / heating source
$SCPe$	equivalent specific cooling power $W/kg_{ads}$	<i>h</i>	hot, heating source
$t$	time, s	<i>n</i>	constant used in eq. 2
$T$	temperature, K	<i>o</i>	outlet
$x$	adsorption uptake, $kg/kg_{ads}$	<i>s</i>	saturation
$x_{eq}$	equilibrium uptake, $kg/kg_{ads}$	<i>w</i>	water

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439

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