Dehumidification Effect of Polymeric Superabsorbent SAP-LiCl Composite Desiccant-Coated Heat Exchanger with Different Cyclic Switching Time

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Abstract: This study investigated a composite polymer desiccant material’s performance, which is prepared by impregnating solid desiccant such as sodium polyacrylate (SAP) on to hygroscopic salts such as lithium chloride (LiCl). Dehumidification performance of the proposed composite polymer desiccant (SAP-LiCl) was analyzed by coating the suitable weight percentage (wt %) of the desiccant onto a single fin-tube heat exchanger (FTHE) system and testing the desiccant-coated heat exchanger (DCHE) in a testing tunnel under various operating conditions. Net dehumidification efficacy of DCHE in terms of sorption and desorption amount and thermal performance (COPth) were analyzed. For instance, with processed air inflow temperature, relative humidity and regeneration temperature setting of 30 °C, 80% RH and 70 °C, DCHE’s sorption, desorption amount and COPth were recorded as high as 945.1 g, 1115.1 g, and 0.39, respectively. It was further realized that the performance of the DCHE could be enhanced by modulating the cyclic switching time for dehumidification and regeneration processes. For instance, with the aforementioned processed airflow conditions, when the cyclic switching time tuned as 60 min instead of 10 min for a total time period of 120 min, there is a net 58% improvement to the COPth of the system. It was further observed that, under the same time period corresponding to the increase in cyclic switching time, the overall COPth can be enhanced; however, the water vapor sorption and desorption amounts of desiccant were decreased.

Keywords: composite polymer desiccant; fin-tube heat exchanger; vapor sorption; vapor desorption; cyclic switching time; COPth

1. Introduction

The last few decades have witnessed rapid urbanization as well as industrialization. It is further predicted that the globe’s primary energy requirement will see at least a 40% surge in the next two decades [1]. It has been further highlighted that 40% of total energy consumption around the world accounted for the buildings from which 50% comes from the air conditioning system [1–3]. It can be noted that 40% of the total air-conditioning load is latent load, which is related to the moisture content handlings of the air-conditioning space [3]. To reduce net latent load, desiccant dehumidifiers have been developed, which is used to remove the moisture from the air either through the absorption or adsorption mechanism. The last few decades have witnessed significant research development towards the design and application of various desiccant dehumidifiers, which can be broadly classified into three major categories, namely, rotary desiccant wheel, fixed bed, and desiccant-coated heat
exchangers (DCHEs). Each technology has their advantages and disadvantages over each other, and extensive details in this regards can be found in several literature studies [1,4–7]. Among all the three aforementioned technologies, DCHEs are considered as the most efficient methodology to reduce the overall cooling load with improved dehumidification with the least energy budget expenditure, hence employed for many real-life applications [5]. Along with air-conditioning technology, DCHEs are being used in absorption chillers, heat pumps [3,8], and even for water harvesting [9].

Desiccant-coated heat exchangers are basically fin-tube heat exchangers (FTHEs), which are comprised of desiccant-coated aluminum (Al) fins, where copper (Cu) tubes are embodied within it [10]. Al fins are coated with either solid or polymer desiccants on it, and when the process air passes through it, the vapors are captured through the desiccants. To achieve isotherm sorption, the cold water is supplied through the copper tubes while ambient air passes through the DCHEs. Meanwhile, hot water is supplied through the copper tubes for the regeneration of captured vapor. It was realized that the efficiency of DCHE largely depends on the desiccant material property and the temperature of working medium supply. Conventional solid desiccant material such as silica gel is widely used desiccant material in DCHE towards moisture removal [3]. To further improve the sorption capability, silica gel was further dispersed within lithium chloride (LiCl), calcium chloride (CaCl₂), and potassium formate (CHKO₂) solution to realize a net improvement in the sorption capability [11,12]. Other desiccant materials such as zeolite, Metal-Organic Frameworks (MOF) and composite carbon desiccant have also shown their efficacy in improving the net isotherm sorption amount of the DCHE [13,14].

To further improve the vapor sorption, material scientists have devised cross-linked polymers with a base of acrylates and acrylamides [15], which are coined as superabsorbent polymers (SAP). SAP possesses superlative capacity towards water absorption as it can retain water up to 2–3 times its original weight [1,16]. At a similar working condition, dehumidification capacity of SAP was found to be superior to other available solid desiccants such as silica gel [15,17]. SAP was further impinged within hygroscopic salts such as LiCl solution to prepare composite super desiccant polymer [16]. The composite super desiccant polymer’s sorption capacity was tested on desiccant wheels, and it was realized that sorption capacity is 2–3 times better than traditional solid desiccant material silica gel [18]. Recent studies have extensively highlighted that the composite desiccant-coated FTHEs demonstrate superlative capacity towards the dehumidification process [7,10]. For instance, in a recent study, Vivek et al. evaluated the PVA (polyvinyl alcohol)-LiCl-coated heat exchanger’s performance that illustrates a 20–60% improvement in the dehumidification capacity [10]. Although the efficacy of the composite polymer SAP-LiCl has shown its efficacy towards improved vapor sorption [16,18], its performance on a FTHE is still unknown. Hence, the objective of this work is to evaluate the dehumidification performance of a SAP-LiCl-coated heat exchanger in various environmental conditions. The appropriate concentration of SAP in the LiCl solution was first determined for efficient vapor sorption and desorption through a static experiment in a controlled programmable constant temperature and humidity chamber. Subsequently, the fin-tube heat exchangers were coated with the optimized amount of the proposed desiccant, and a dynamic experiment within an air tunnel was carried out to evaluate the performance of the proposed composite desiccant in various process airflow conditions and cycling switching times between the dehumidification and regeneration process.

2. Materials and Experimental Methods

2.1. Desiccant Coating Preparation and Static Performance Measurement

To prepare the composite desiccant solution, appropriate amounts of lithium chloride (LiCl) were mixed with water maintained at a temperature of 50 °C. Subsequently, approximate amounts of superabsorbent polymers (SAP) were added into the aqueous LiCl solution and constantly stirred until a desired homogeneous solution was obtained. Three resultant desiccant solutions with three different concentrations of SAP (16.6 wt %, 33.3 wt %, and 50.0 wt %) were tested to find out the efficacy of the composite desiccant polymer material and its optimal concentration towards moisture
sorption and desorption. The resultant solution uniformly adhered to an aluminum sheet of 4 × 4 cm (width × breadth) dimension. Subsequently, the desiccant coated on the aluminum sheet underwent a baking process in an oven at a temperature of 80 °C for 5 h towards the hardening and drying of the desiccant materials. The vapor sorption and desorption capacity of the desiccant-coated Al sheets were quantified for five different temperature settings in a controlled programmable constant temperature and humidity chamber. The timeframes of both the tests were set to a total period of 15,000 s, and the moisture sorption and desorption were recorded at every 1000 s, by measuring the weight of the aluminum sheet.

2.2. Dynamic Performance Measurement

2.2.1. Immersion Coating on FTHE

To remove any unnecessary pollutants from the surface of the heat exchanger fins, it was first immersed in a mixture of citric acid and water solution and underwent an ultrasonic bath at a temperature of 50 °C for a time period of 90 min. Subsequently, the heat exchanger was further cleaned with water and air-dried. The optimized concentration of desiccant material was further uniformly coated on the heat exchanger through immersion coating technology, and the whole setup underwent baking in an industrial oven maintained at a temperature of 80 °C for a period of 60 h.

2.2.2. Experimental Setup

The schematic illustration of the experimental setup is illustrated below in Figure 1. The experimental setup was comprised of an acrylic chamber, an air environment-controlled chamber, two water baths, ultrasonic humidifiers, fans, and a desiccant-coated fin-tube heat exchanger. During the experiment, the air was supplied from the environment-controlled chamber at a flow rate of 1.8 m³/min with constant temperature and humidity condition. Aluminum honeycomb panels were fitted on the front and backend of the DCHE chamber to supply a uniform airflow. Six temperature and humidity sensors were fitted before and after DCHE to record the incoming and processed airflow. For the regeneration process, the water was supplied to the DCHE chamber through a water pump, and the temperature was constantly monitored through a temperature sensor.

![Figure 1. Schematic illustration of the experimental setup.](image-url)
2.2.3. Theoretical Performance Analysis

The moisture removal rate \( (M_{vap}) \) of the DCHE can be quantified by measuring the inlet and outlet humidity ratios and the mass flow rate of the process air as provided in the underneath Equation (1).

\[
M_{vap} = \dot{m}_a (W_{a, \text{out}} - W_{a, \text{in}}), \tag{1}
\]

where \( \dot{m}_a \) is the mass flow rate of air and \( W_{a, \text{in}} \) and \( W_{a, \text{out}} \) are the inlet and outlet humidity ratios of the process air, respectively.

The vapor sorption or desorption amounts \( (G_{vap}) \) were determined through Equation (2) by taking integral of the moisture removal rate with respect to time \( (\tau) \).

\[
G_{vap} = \int_0^\tau |M_{vap}| \, dt, \tag{2}
\]

where \( \tau \) is the time period for the vapor sorption or desorption process.

The vapor sorption or desorption amount \( (G_a) \) of desiccant material per unit area \( (\text{g/cm}^2) \) can be further quantified by dividing the vapor sorption or desorption amounts \( (G_{vap}) \) by the area of fin \( (A_{\text{fin}}) \),

\[
G_a = \frac{G_{vap}}{A_{\text{fin}}}. \tag{3}
\]

The vapor sorption or desorption ability per unit mass of the desiccant \( (G_m) \) was further determined by dividing vapor sorption or desorption amounts \( (G_{vap}) \) by the mass of the desiccant \( (M_{\text{des}}) \) coated on DCHE,

\[
G_m = \frac{G_{vap}}{M_{\text{des}}}. \tag{4}
\]

To find out the overall energy efficiency of DCHE, the average thermal performance \( \text{COP}_{th} \) was quantified, which is defined as the ratio of the cooling capacity of air \( (Q_{\text{cool}}) \) to the heat exchange of water during an effective dehumidification \( (Q_{\text{deh}}) \) and regeneration process \( (Q_{\text{reg}}) \).

\[
\text{COP}_{th} = \frac{Q_{\text{cool}}}{Q_{\text{deh}} + Q_{\text{reg}}} = \frac{\int_0^{\tau_a} \dot{m}_a \times \Delta H \, dt}{\int_0^{\tau_a+\tau_r} \left\{ (\dot{m}_w \times C_p \times \Delta T_{\text{deh}}) + (\dot{m}_w \times C_p \times \Delta T_{\text{reg}}) \right\} \, dt} \tag{5}
\]

where \( \dot{m}_a \) and \( \dot{m}_{\text{water}} \) are the mass flow rate of the process air and the regeneration hot water, respectively, in kg/s, \( \Delta H \) represents the difference in enthalpies of the process before and after DCHE, respectively, in kJ/kg, and \( C_p \) is the specific heat of the regeneration hot water at a constant pressure in kJ/kg K. \( \Delta T \) is the temperature difference of water supply to \( (T_{w, \text{in}}) \) and exhausting \( (T_{w, \text{out}}) \) from the DCHE, respectively. \( \tau_a \) and \( \tau_r \) are the time periods for the vapor sorption and desorption process.

2.2.4. Uncertainty Analysis

The system uncertainties may emanate from instruments, physical environment, and human inadequacies. The system uncertainties for this work were quantified using the previously established protocols [19,20]. The instrument errors that influence the uncertainty analysis majorly emanate from the programmable constant temperature and humidity system (Model-150-00-SP-SD, Giant Force Instrument Enterprise Co., Ltd., Taiwan, 0–100 °C ± 0.3 °C, 10–98% RH ± 3% RH), temperature-controlled water bath (Model-D-620, Great Tide Instrument Co., Ltd., Taiwan, −20–100 °C ± 0.1 °C), precision electronic scale (Model-AB323, Dehe Weighing Apparatus Co., Ltd., Taiwan, 0–320 g ± 0.003 g), electric heating plate (Model-HP303D, Ten Billion Technology Co., Ltd., Taiwan, 0–350 °C ± 1 °C), anemometer (Model-TES-3142, TES Co., Ltd., Taiwan, 1–80 m/s ± 3%), temperature probe (Model-HF532, Hsing Nan Import & Export Co., Ltd., Taiwan, −40–85 °C ±
0.1 °C), humidity probe (Model-HC2-IC302, Hsing Nan Import & Export Co., Ltd., Taiwan, 0–100% RH ± 0.8% RH) and liquid flow meter (Model-NW20-NTN, JETEC Electronics Co., Ltd., Taiwan, 1.5–20 L/min ± 2%). The uncertainty in moisture removal rate ($M_{vap}$), vapor sorption or desorption amounts ($G_{vap}$), vapor sorption or desorption ability per unit area ($G_a$), vapor sorption or desorption ability per unit mass ($G_m$), COP$_{th}$ and energy consumption (E) were quantified within the range of ±2.2%, ±2.2%, ±4.2%, ±6.5%, ±3.7%, and ±4.75%, respectively. An extensive detail regarding the uncertainty analysis can be found in one of our recently published articles [20].

3. Results and Discussions

3.1. Performance Analysis of Desiccant in a Static Test Environment

To test the weight concentration of sodium polyacrylate (SAP) in the composite desiccant solution’s (SAP-LiCl) sorption amount, experiments were conducted with a single aluminum sheet coated with 16.6 wt %, 33.3 wt %, and 50.0 wt % of SAP in the desiccant solution, and the experiment was conducted in a programmable constant temperature and humidity chamber by maintaining the humidity ratio at 80% RH. The sorption kinetics was recorded at a different temperature of 20 °C, 25 °C, 30 °C, 35 °C, and 40 °C for a time period of 15,000 s (left column of underneath Figure 2). It was observed that, irrespective of SAP concentration (i.e., 16.6 wt %, 33.3 wt %, and 50.0 wt %), the sorption amount of the desiccants increases corresponding to the increase in temperature. For instance, after 15,000 s, the sorption amount for a 16.6 wt % desiccant-coated Al sheet was measured as 2.005 g/g when the chamber temperature was maintained at 20 °C. With a similar desiccant concentration, the sorption amount increases to 3.916 g/g, when the chamber temperature was maintained at 40 °C, suggesting a 48% net increment in the sorption amount. A similar trend in the vapor sorption was recorded for the other two weight concentrations (i.e., 33.3 wt % and 50.0 wt %). Upon comparison, it was found that, under similar working concentrations, the proposed SAP-LiCl-coated FTHE performs better in terms of vapor sorption compared to its currently available other counterparts. For instance, the maximum sorption amount recorded for PVA-LiCl-coated heat exchanger was accounted as 0.8 g/g, when the chamber temperature was maintained at 30 °C with 80% RH [10]. At similar working conditions, the net sorption amount recorded for SAP-LiCl-coated FTHE was recorded as high as 1.8 g/g, suggesting a 125% increment in the sorption amount. The improvement in DCHE’s performance can be explained by shedding light on the molecular structure of the composite desiccant. In particular, compared to other solid desiccants, SAP has a smaller surface area. However, when impregnated with salt such as lithium chloride (LiCl), the surface area of polymer desiccant enhances due to salt crystal formation, which further enhances its vapor sorption capability [16]. Furthermore, upon quantification, the vapor sorption rate was found to attain equilibrium in quick succession of 6000–8000 s. at a relatively lower temperature of 20 °C, 25 °C, and 30 °C for 16.6 wt % and 33.3 wt % (ESI Figure S1).

To evaluate the vapor desorption capability of the desiccant during the regeneration process, experiments were conducted at a temperature of 40 °C, 50 °C, 60 °C, 70 °C, and 80 °C by maintaining the relative humidity at 80% RH with processed air inflow of 30 °C. It was observed that the Al sheet with 16.6 wt % yields a poor desorption irrespective to change in temperatures. For 33.3 wt % and 50.0 wt %, vapor desorption kinetics significantly improves corresponding to an increase in the temperature. At relatively lower temperatures such as 40 °C and 50 °C, desiccants cannot be completely dehydrated for all three desiccant concentrations. However, at a higher desiccant concentration (33.3 wt % and 50.0 wt %), vapor desorption significantly improves when regeneration temperature was set at 70 °C and 80 °C as the desiccant-coated aluminum sheets can be completely dehydrated to zero. The superior desorption characteristics of the proposed desiccant at higher temperatures might be due to its large vapor capacity during the dehumidification process, which requires higher regeneration temperature towards its complete dehydration. Further, considering that the 16.6 wt % desiccant-coated Al sheets possess a poor desorption and 33.3 wt % and 50.0 wt % desiccant-coated Al sheets showed a similar
trend in sorption as well as desorption, 33.3 wt % was selected as the final coating concentration to apply on the fin-tube heat exchanger for dynamic testing operation.

![Figure 2](image)

**Figure 2.** Vapor sorption and desorption kinetics of SAP-LiCl desiccant at the various operating conditions. The first row (a,b), second row (c,d), and third row (e,f) illustrate the vapor sorption and desorption capability of the proposed composite desiccant with 16.6 wt %, 33.3 wt %, and 50 wt %, respectively. The results are depicted as Mean ± SD (n = 4).

3.2. Performance Analysis of Composite Desiccant-Coated FTHEs in a Dynamic Test Condition

3.2.1. Moisture Removal Capability of Desiccant-Coated FTHEs at Different Humidity Conditions

The underneath Figure 3 illustrates the distribution curve of the air humidity ratio of the desiccant-coated heat exchanger with selected concentration of 33.3 wt % under different process air humidity conditions, with a processed air inflow maintained at 30 °C. During the experiment, cold and hot water alternatively supplied to the heat exchanger towards dehumidification as well...
as regeneration processes with process times of 30 min for each cycle. It can be noted that, during the dehumidification process, the cold water is supplied at 25 °C, whereas, during the regeneration process, the supplied water temperature was maintained at 50 °C. As observed, with a relatively lower inlet air humidity condition of 60% RH, a relatively small change in the humidity ratio was observed during the dehumidification and regeneration process. However, under high humidity conditions of 70% RH and 80% RH, a significant change in the humidity ratio was observed during both the dehumidification as well as the regeneration process. For all the three test conditions, approximately 7–9 min were required to attain the saturated state. It was further observed that the humidity condition of inflow air significantly affects the time required to achieve regeneration. For instance, with 60% RH, the regeneration was complete after 6 min, whereas it approximately takes 8 min when the humidity of air was maintained at 80% RH. However, at this point, it is not very clearly known how the sorption and desorption amount of proposed DCHE changes under various operating conditions such as different regeneration temperatures, process air inflow temperature, switching cyclic time period between humidification and regeneration processes, etc. To elucidate it further, experiments were conducted and illustrated in the following sections.

![Distribution curves of the outlet air humidity ratio of SAP-LiCl-coated heat exchangers at various humidity conditions of 60% RH, 70% RH and 80% RH.](image)

**Figure 3.** Distribution curves of the outlet air humidity ratio of SAP-LiCl-coated heat exchangers at various humidity conditions of 60% RH, 70% RH and 80% RH. The processed air inflow was maintained at 30 °C and the water temperature was maintained at 25 °C and 50 °C during the dehumidification and regeneration process.

3.2.2. Vapor Sorption or Desorption Capability of Desiccant-Coated FTHEs at Different Operating Conditions

Tables 1 and 2 illustrate the vapor sorption and desorption amount ($G_{vap}$), sorption and desorption amount per unit area ($G_a$), sorption and desorption amount per unit mass ($G_m$) of the composite SAP-LiCl-coated FTHE under different regeneration temperatures (50 °C, 60 °C, and 70 °C) and humidity conditions (60% RH, 70% RH, 80% RH) with two processing air inflow temperature settings of 25 °C and 30 °C were tested. To avoid terminological confusion, the air inflow temperature settings of 25 °C and 30 °C were coined as Case-I and Case-II, respectively. The vapor sorption and desorption abilities were noticed to be increased, corresponding to the increases in the regeneration temperature and relative humidity for both Case-I and Case-II. It was further observed that at a higher processing air
inflow temperature of 30 °C (Case-II), vapor sorption and desorption abilities of the SAP-LiCl-coated heat exchanger improves significantly. For instance, with a regeneration temperature setting at 70 °C with a humidity air ratio of 80% RH, the sorption amount ($G_{vap}$) of the DCHE was accounted as 945.06 g and 1115.1 g, respectively. This suggests a net 17% improvement in the net sorption ability of the DCHE at a higher processing air inflow temperature of 30 °C (Case-II). A similar trend in the improvement was further noticed for the DCHE’s sorption and desorption amount per unit area ($G_a$), sorption and desorption amount per unit mass ($G_m$) for all the other test conditions. This improvement might result from the increase in the humidity ratio of the process air due to the increase in the dry bulb temperature under the same relative humidity (% RH) in the processed air. Upon comparison, it was further observed that, under a similar working condition, the sorption and desorption amount of SAP-LiCl-coated FTHE was found to be better compared to silica gel-coated FTHE. For instance, at a processing inlet airflow temperature setting of 30 °C with humidity ratio of 70% RH, the sorption amount ($G_{vap}$) of the silica gel-coated FTHE was accounted as low as 86.75 g upon compared to a sorption amount of 1115.1 g obtained for an SAP-LiCl-coated FTHE [15]. A similar trend in the improvement was observed for all the remaining test conditions. Hence it can be concluded that the proposed composite desiccant possesses a superlative capability towards moisture removal in any weather condition.

### Table 1. Vapor sorption and desorption abilities of SAP-LiCl-coated FTHEs at different operation conditions with processed inlet airflow temperature settings of 25 °C (Case-I) and 30 °C (Case-II).

| Regeneration Temperature | Sorption Amount | Desorption Amount | Sorption Amount | Desorption Amount | Sorption Amount | Desorption Amount |
|--------------------------|-----------------|-------------------|-----------------|-------------------|-----------------|-------------------|
|                          | 60% RH          | 70% RH            | 80% RH          | Case-I—Processed air inflow temperature of 25 °C |
| $G_{vap}$ (g)            |                 |                   |                 | 50 °C              | 252.3           | 361.8             | 289.6             | 424.9             | 348.9             | 457.8             |
| 50 °C                    | 287.7           | 526.6             | 346.7           | 597.4             | 402.2           | 709.1             |
| 70 °C                    | 315.0           | 667.6             | 380.3           | 777.6             | 442.5           | 945.1             |
| $G_a$ (g/cm²)            |                 |                   |                 |                   | 50 °C            | 0.052             | 0.075             | 0.060             | 0.088             | 0.072             | 0.095             |
| 50 °C                    | 0.060           | 0.109             | 0.072           | 0.124             | 0.083           | 0.147             |
| 70 °C                    | 0.065           | 0.138             | 0.079           | 0.161             | 0.092           | 0.196             |
| $G_m$ (g/g)              |                 |                   |                 |                   | 50 °C            | 0.661             | 0.948             | 0.759             | 1.114             | 0.914             | 1.200             |
| 50 °C                    | 0.754           | 1.380             | 0.909           | 1.566             | 1.054           | 1.859             |
| 70 °C                    | 0.826           | 1.750             | 0.997           | 2.038             | 1.160           | 2.478             |
|                          |                 |                   |                 | Case-II—Processed air inflow temperature of 30 °C |
| $G_{vap}$ (g)            | 50 °C            | 282.9             | 420.5           | 309.0             | 504.0           | 342.0             | 584.4             |
| 50 °C                    | 303.7           | 641.3             | 396.0           | 732.9             | 418.0           | 1006.1            |
| 70 °C                    | 336.8           | 800.9             | 432.9           | 967.9             | 486.4           | 1115.1            |
| $G_a$ (g/cm²)            |                 |                   | 50 °C            | 0.059             | 0.087           | 0.064             | 0.104             | 0.071             | 0.121             |
| 50 °C                    | 0.063           | 0.133             | 0.082           | 0.152             | 0.087           | 0.208             |
| 70 °C                    | 0.070           | 0.166             | 0.090           | 0.201             | 0.101           | 0.231             |
| $G_m$ (g/g)              | 50 °C            | 0.742             | 1.102           | 0.810             | 1.321           | 0.879             | 1.502             |
| 50 °C                    | 0.796           | 1.681             | 1.038           | 1.921             | 1.075           | 2.586             |
| 70 °C                    | 0.883           | 2.099             | 1.135           | 2.542             | 1.250           | 2.867             |
Table 2. Vapor sorption and desorption amount ($G_{vap}$) of SAP-LiCl coated for the first and overall cycles with different inlet airflow conditions (Case-I, Case-II, and Case-III).

| Case   | Dehumidification and Regeneration Amount ($G_{vap}$) for an Individual First Cycle and Overall Cycle Time. |
|--------|----------------------------------------------------------------------------------------------------------|
| Cycle  | First Cycle | Overall | First Cycle | Overall | First Cycle | Overall | First Cycle | Overall | First Cycle | Overall | First Cycle | Overall | First Cycle | Overall |
| Time (min) | Sorption Amount (g) | Desorption Amount (g) | Sorption Amount (g) | Desorption Amount (g) | Sorption Amount (g) | Desorption Amount (g) |
| 10     | 67          | 801     | 86          | 1026     | 85         | 1018     | 101         | 1203     | 100         | 128      | 1204      | 1536     |
| 20     | 127         | 764     | 186         | 1113     | 161        | 964      | 243         | 1456     | 198         | 280      | 1192      | 1680     |
| 40     | 222         | 666     | 311         | 933      | 287        | 861      | 401         | 1203     | 361         | 495      | 1084      | 1486     |
| 60     | 308         | 616     | 398         | 796      | 386        | 772      | 538         | 1076     | 504         | 683      | 1009      | 1366     |

3.2.3. Thermal Coefficient of Performance ($\text{COP}_{th}$)

The overall thermal coefficients of performance ($\text{COP}_{th}$) of the proposed DCHE under different operating conditions were quantified and illustrated in Figure 4 below for both the test cases (processed inlet airflow temperature setting of 25 °C (Case-I) and 30 °C (Case-II)). It was observed that the value $\text{COP}_{th}$ is directly proportional to the relative humidity in the process air for both the test cases. The overall $\text{COP}_{th}$ significantly improved when the air inflow temperature was set at 30 °C (Case-II) rather than 25 °C (Case-I). For instance, with a regeneration temperature setting at 70 °C with 80% RH, the $\text{COP}_{th}$ of DCHE was found to be a maximum of 0.39 and 0.34, respectively, for both Case-II and Case-I, suggesting a 17% net improvement. As mentioned in the earlier section, corresponding to the increase in the dry bulb temperature under the same relative humidity of the process air increases the vapor pressure in process air that boosts sorption abilities resulting in a net enhancement of the system’s overall $\text{COP}_{th}$. It can be further noted that, with a similar environmental setting, with a processed airflow temperature setting of 30 °C, the $\text{COP}_{th}$ of MOF-coated heat exchanger has a value of 0.1662 [ESI Table S1, unpublished work]. Moreover, the overall $\text{COP}_{th}$ of the DCHE was found to be inversely proportional to the increase in regeneration temperature. For instance, in test Case-II, with humidity ratio of 80% RH, the $\text{COP}_{th}$ was quantified as 0.39, 0.33, and 0.31 when the regeneration temperature was set at 50 °C, 60 °C, and 70 °C, respectively. The results above further explain that the proposed composite polymer (SAP-LiCl)-coated heat exchanger depicts a net 60% improvement in the overall $\text{COP}_{th}$ compared to the MOF-coated heat exchanger.
Figure 4. Overall thermal performance coefficient ($\text{COP}_{th}$) of SAP-LiCl composite polymer desiccant under different regeneration temperature and humidity conditions at processed air inflow temperature setting of (a) 25 °C (Case-I) and (b) 30 °C (Case-II), respectively. The uncertainty of $\text{COP}_{th}$ was found within a range of ±3.7%.
3.2.4. Moisture Removal Capability of Desiccant-Coated FTHE with Different Cyclic Switching Time

It has been earlier observed that, with suitable switching time in the dehumidification and regeneration process, the performance of the DCHE can be improved significantly \[10,15\]. To evaluate the performance of SAP-LiCl-coated FTHEs under humidity conditions of 70% RH and regeneration temperature of 50 °C, an experiment was conducted by tuning to various switching times of 10 min, 20 min, 40 min, and 60 min for an overall time period of 120 min. The tests were conducted for three processes air inflow temperature settings of 25 °C (Case-I), 30 °C (Case-II) and 35 °C (Case-III). The humidity ratio curves for the three test cases are illustrated in Figure 5 below. It was observed for all the tested cyclic switching times that the magnitude of humidity ratio was higher for Case-III compared to Case-II and Case-I. This describes even with the tuning of switching time that the DCHE performs better in terms of vapor sorption and desorption, with a higher processed air inflow temperature setting. It was further observed that, corresponding to the increase in cyclic switching time, the sorption and desorption amounts during the dehumidification and regeneration process increases significantly during the first individual cycle (Table 2). This resulted from the longer sorption and desorption time periods in the case with a longer switching time cycle. However, the net sorption as well as desorption amount for an overall cycle decreases compared to a shorter cyclic switching time after an overall time period of 120 min. For instance, in test Case-II, the vapor desorption amount for an individual and overall cycle was accounted as 101 g and 1203 g with a cyclic switching time of 10 min, which further changes to 538 g and 1076 g, respectively, when the cyclic switching time was tuned to 60 min, suggesting an overall decrease in the sorption and desorption amounts with a larger cyclic switching time. For the test case with a shorter switching cycle time, even the sorption and desorption amount in the first individual cycle is less, yet it significantly increases corresponding to the overall time period due to the increase in number of cycles. Similarly, for a longer switching cyclic time, although the vapor sorption and desorption capabilities (g/g) are greater in the first switching cycle, as the system underwent less individual cycle, the net vapor sorption and desorption amounts decrease. For instance, for Case-I, the net vapor sorption amount for the first cycle was quantified as 67 g for a 10 min switching time cycle, which further improves to 308 g when the cyclic switching time was tuned to 60 min, suggesting a net improvement in vapor sorption amount corresponding to the first cycle corresponding to an increase in switching cycle time. However, the system underwent 12 and 2 individual cycles for an overall time period of 120 min, when switching cycle time is tuned as 10 min and 60 min, respectively. Corresponding to 12 and 2 individual cycles, the overall sorption amount was recorded as 801 g and 616 g after the overall cycle of 120 min. Similar trends in results are illustrated for sorption and desorption amount for the test cases I, II, and III, suggesting that a larger cyclic switching time may reduce the vapor sorption and desorption amount for an overall cycle. However, from the above results, it was not conclusive how the overall performance of the DCHE was affected in terms of expenditure. To correlate the energy budget to the DCHE’s performance corresponding to the tuning of cyclic switching time, the thermal coefficient performance (COP_{th}) of the DCHE corresponding to different cyclic switching time was quantified as described in the next section.
Figure 5. Distribution curves of the outlet air humidity ratio corresponding to cyclic switching times of 10 min, 20 min, 40 min, and 60 min for DCHE under similar regeneration temperature of 50 °C and humidity conditions (70% RH) at processed air inflow temperature settings of (a) 25 °C (Case-I), (b) 30 °C (Case-II), and (c) 35 °C (Case-III), respectively.
3.2.5. The Overall and Instantaneous COP\textsubscript{th} of Desiccant-Coated FTHE Corresponding to Different Cyclic Switching Times

The overall coefficient of performance (COP\textsubscript{th}) of SAP-LiCl-coated FTHE under different cyclic switching times of 10 min, 20 min, 40 min, and 60 min for the first individual cycle and for an overall time period of 120 min were quantified and illustrated in Table 3 below. It was observed that, irrespective of the cyclic switching time, the individual (first cycle) as well as overall COP\textsubscript{th} was highest for test Case-III. For instance, at inlet airflow temperature of 25 °C (Case-I), the overall COP\textsubscript{th} for the four different cyclic switching times were accounted as 0.1940, 0.2673, 0.3084, and 0.3254, respectively which, further improved to 0.3693, 0.4939, 0.5913, and 0.6207, respectively, when the inlet airflow was tuned to 35 °C (Case-III). This suggests a net improvement in the thermal coefficient, with an increment in the processed airflow temperature. It was further observed that, irrespective of inflow condition, the COP\textsubscript{th} value increases, corresponding to an increase in the cyclic switching time. For Case-III, during the first cycle, COP\textsubscript{th} was accounted as 0.3612, 0.4887, 0.5895, and 0.6318, respectively, when cyclic switching time was set as 10 min, 20 min, 40 min, and 60 min, respectively, suggesting an increment in the COP\textsubscript{th} with larger cyclic switching times. For instance, with processed air inflow temperature, relative humidity and regeneration temperature setting of 30 °C, 70% RH and 70 °C, there was a net 58% improvement in the COP\textsubscript{th} when the cyclic switching time was tuned to 60 min instead of 10 min. Similarly, with analogous inlet process airflow condition, when the inlet air temperature was set at 35 °C, there was a net 68% improvement. A similar trend of increment in COP\textsubscript{th} was noticed for all the test cases for an overall cyclic time period of 120 min, suggesting an improvement in energy efficiency corresponding to larger cyclic switching times. However, as delineated in the previous Section 3.2.4 (Table 2), a longer cyclic switching time reduces the vapor sorption as well as desorption amounts for an overall time period of 120 min. However, it was observed hereby that there is an overall improvement to the COP\textsubscript{th} corresponding to longer cyclic switching times. To shed light on this aspect and to delineate the underlying phenomena, the energy expenditure in terms of cooling capacity during the sorption process as well as the energy consumption in sorption and desorption processes for Case-III (maximum improvement in the COP\textsubscript{th}) were quantified for the first individual cycle and illustrated in Figure 6 below. As observed, corresponding to an increase in the cyclic switching time, the cooling capacity in the sorption process and the energy consumption in both processes gradually increase as well. However, with the increase in the switching cyclic time, the increment of energy consumption in both processes is less than that of the cooling capacity in the sorption process. This indicates that, in the case of a short switching cyclic time, the energy loss in terms of the cooling capacity gain is greater due to frequent operational mode switching, which results in the COP\textsubscript{th} value in the individual cycle gradually increasing with the increase in the switching cyclic time. The improvement in each individual cycle further improves the overall COP\textsubscript{th} in an overall cycle.

**Table 3.** COP\textsubscript{th} of SAP-LiCl-coated FTHE under different cyclic switching times for both test cases.

| Cycle Time (min) | First Cycle | Overall | First Cycle | Overall | First Cycle | Overall |
|------------------|-------------|---------|-------------|---------|-------------|---------|
| 10               | 0.1890      | 0.1940  | 0.2799      | 0.2885  | 0.3612      | 0.3693  |
| 20               | 0.2649      | 0.2673  | 0.3838      | 0.3837  | 0.4887      | 0.4939  |
| 40               | 0.3033      | 0.3084  | 0.4539      | 0.4659  | 0.5895      | 0.5913  |
| 60               | 0.3213      | 0.3254  | 0.4561      | 0.4557  | 0.6318      | 0.6207  |
The dehumidification ability of SAP-LiCl-coated FTHE was analyzed under different switching time cycles, and it was observed that with the increase in the switching time cycle, the overall thermal coefficient of performance increases corresponding to the decrease in the overall sorption and desorption ability of the DCHEs. Energy analysis suggests that a longer switching cycle decreases energy loss during each cycle, resulting in a net improvement in the thermal coefficient of performance ($\text{COP}_{th}$) of DCHE.

4. Conclusions

In this study, a composite polymer desiccant material, SAP-LiCl, was tested for its moisture removal efficacy through static and dynamic testing conditions. The results illustrate that the proposed desiccant possesses a superlative dehumidification capacity compared to the other currently available desiccant materials. Based on the experimental results, the following conclusions can be derived as listed underneath.

1. The static experiment was carried out with a desiccant-coated Al sheet in a programmable constant temperature and humidity system. For superior sorption as well as desorption functionality, the final weight concentration (wt %) of SAP in the polymer composite desiccant solution (SAP-LiCl) was optimized as 33 wt %.

2. Under high humidity conditions (70% RH and 80% RH), a significant change in the humidity ratio curve was noticed during both the dehumidification as well as the regeneration process, suggesting the efficacy of the proposed desiccant-coated FTHE in high humidity conditions.

3. The sorption and desorption amount of SAP-LiCl-coated FTHE was found to be superior compared to silica gel-coated FTHE. For instance, at a processing inlet airflow temperature setting of 30 °C with humidity ratio of 70% RH, the sorption amount ($G_{\text{vap}}$) of the SAP-LiCl-coated FTHE was accounted as 1115.1 g compared to a value of 86.75 g of silica gel-coated FTHE.

4. The sorption amount and $\text{COP}_{th}$ of DCHE were found to be better with processed air inflow with a temperature setting of 30 °C compared to 25 °C. With processed air inflow temperature, relative humidity and regeneration temperature setting of 30 °C, 80% RH and 70 °C, DCHE’s $\text{COP}_{th}$ is recorded as high as 0.39.

5. The dehumidification ability of SAP-LiCl-coated FTHE was analyzed under different switching time cycles, and it was observed that with the increase in the switching time cycle, the overall thermal coefficient of performance increases corresponding to the decrease in the overall sorption and desorption ability of the DCHEs. Energy analysis suggests that a longer switching cycle decreases energy loss during each cycle, resulting in a net improvement in the thermal coefficient of performance ($\text{COP}_{th}$) of DCHE.
These proposed desiccant’s applications can also be extended to absorption chillers, heat pumps, and water regenerators and is the future direction of our ongoing research.

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Nomenclatures

- \( M_{vap} \) Moisture removal rate
- \( m_a \) Mass flow rate of air
- \( W_{a,in} \) Humidity ratio of process air inlet
- \( W_{a,out} \) Humidity ratio of process air outlet
- \( G_{vap} \) Vapor sorption/desorption amount
- \( T \) Period for vapor sorption or desorption
- \( G_a \) Vapor sorption/desorption amounts per unit area of desiccant
- \( A_{fin} \) Total area of the fins
- \( C_m \) Sorption or desorption ability per unit mass of the desiccant
- \( COP_{th} \) Coefficient of thermal performance
- \( C_P \) Specific heat
- \( Q_{reg} \) Average heat exchange of water in an effective regeneration process
- \( Q_{cool} \) The ratio of the total energy exchange of the processed air in an effective dehumidification process
- \( RH\% \) Relative humidity in percentage
- \( T_{h,in} \) Temperatures of water supply to DCHE
- \( h_{a,in} \) Enthalpies of the process air after of DCHE
- \( h_{a,out} \) Enthalpies of the process before of DCHE
- \( m_{water} \) Mass flow rate of water
- \( T_{h,out} \) Temperatures of water exhaust from DCHE

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