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1. Introduction

Biomass is a promising clean energy source that has the potential to replace crude oil. The net CO₂ emissions caused by burning biomass are theoretically significantly less than those produced from burning fossil fuels. This is due to the ability of biomass to reabsorb the emitted CO₂ from the atmosphere as it grows during the photosynthesis process.¹ Biomass usage for energy dates back to when burning wood was used for heating and cooking.² Later on, other processes were developed to convert biomass into an energy crop including biochemical and thermochemical techniques. Conventional biomass used in such processes includes sugar, starches and vegetable oils. More recently, research has shifted to algae and cellulosic/ lignocellulosic biomass species.²

Pyrolysis is a single step thermochemical process that uses heat in the absence of O₂ to decompose biomass into liquid (bio-oil), solid (biochar) and gaseous products.³ A variety of lignocellulosic-based biomass species have been explored for their potential to produce fuel and specialty chemicals.⁴ An interesting plant that is gaining more attention recently is *Salicornia bigelovii* (SB). SB is a plant that belongs to the halophyte species. Halophytes are salt and heat tolerant plants that can grow under harsh conditions.⁵ Therefore, it is possible to grow SB on marginal soils and irrigate with saline water. SB can be

Interactions in co-pyrolysis of *Salicornia bigelovii* and heavy fuel oil[†]

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The interactions in the co-pyrolysis of biomass from *Salicornia bigelovii* (SB) with heavy fuel oil (HFO) were studied through an investigation of the yields, kinetics, and the quality of bio-oil and biochar produced. Co-pyrolysis experiments were performed at 550 °C in a quartz tube with three samples of different ratios of SB and HFO. The decomposition of the samples was investigated using a thermogravimetric analyzer (TGA). The effect of changing the geometry of the sample holder was also investigated in the TGA. The addition of HFO enhanced the volatilization of the sample which resulted in a decrease in char yield and an increase in gas yield. Mixing also increased reactivity and lowered the apparent activation energies in the last stage (227–500 °C) of the co-pyrolysis process. The amount of oxygenated compounds in the bio-oil collected from the mixtures was reduced. The selectivity towards aliphatic and aromatic hydrocarbons increased with an increased addition of HFO and the selectivity towards acids and nitrogen containing groups decreased. Co-pyrolysis of SB and HFO has the potential to produce better quality biofuels through an environmentally friendly process.

directly converted to fuel *via* pyrolysis and has been studied in different environments.⁶⁻⁸ The pyrolysis products of other halophytes such as *Tamarix chinensis*, *Suaeda salsa*, and *Phragmites australis* have also been explored.^{9,10}

Pyrolysis bio-oil has potential in becoming a great candidate for energy generation. However, bio-oil still falls short when compared to fossil fuels due to the presence of heteroatoms, namely, N, O and S. Thus far, bio-oil has been used as a lowgrade fuel,¹¹ a source of value added chemicals,¹² a binder¹³ and a functional carbon material.^{14,15} However, the use of bio-oil as transportation fuel is challenging due to its high content of oxygen and water, among other properties.¹⁶ This leads to issues of corrosion, chemical stability and low calorific value during the storage and use of bio-oil.^{17,18} Therefore, bio-oil processing is needed and multiple techniques have been developed in the past to treat bio-oil. Some examples of such techniques are: hydrotreating, hydrocracking, and solvent addition.¹⁹

A relatively newer technique to enhance the quality of bio-oil is co-pyrolysis, which is the simultaneous pyrolysis of two or more materials without requiring an additional reaction vessel, solvent, or catalyst, although a catalyst may be used to further improve the bio-oil quality.^{17,20} Co-pyrolysis of biomass with plastics has been extensively studied by numerous researchers.^{21–25} These studies have shown that co-pyrolysis of mixtures results in enhanced conversion of biomass to volatiles and gases resulting in lower char yields.^{22–24} The collected char from co-pyrolysis of pine cone with synthetic polymers had a higher calorific value compared to char from biomass pyrolysis alone.²⁵ The bio-oil from co-pyrolysis of biomass and highdensity polyethylene had 76% lower viscosity and lower water

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and oxygen content.²⁶ Studies report that plastics contain large amounts of hydrogen and almost no oxygen, therefore could adjust the content of hydrogen, carbon, and oxygen in the biooil produced.²⁵⁻²⁷ Under co-pyrolysis, scientists speculate that the oxygen in the biomass reacts with the hydrogen and hydrogen radicals from plastics to produce $H_2O.^{28,29}$ This modifies the oxygen content and results in a more stable, less corrosive bio-oil.

Another rich source of hydrogen that can be used in biomass co-pyrolysis is heavy fuel oil (HFO). HFO is a black or brown viscous liquid with a tar-like consistency. It is a mixture of the residue of vacuum distillation and light distillate (cutter stock), that is difficult to burn because of its high concentration of complex hydrocarbons, known as asphaltene, that can cause fouling and corrosion in engines.^{30,31} HFO is mainly used in marine transportation, and although it is considered a low quality transportation fuel, 95% of marine engines use it due to its low cost.32 HFO, like plastics, is abundant in hydrogen and deficient in oxygen relative to biomass. Catalytic and noncatalytic gasification of HFO has been widely studied for the production of clean fuels.33 However, there are limited studies that have reported on the use of heavy fuels as feedstock or cofeedstock in pyrolysis, although petroleum/oily sludge has been used as a co-feed in biomass pyrolysis. The pyrolysis of mixtures of oily sludge with different biomass, such as sawdust, walnut shell, rice husk, and apricot shell, resulted in enhancements in heating value and H/C ratio.34,35 Moreover, bio-oil collected from co-pyrolysis of rice husk with oily sludge contained fewer oxygenates and a higher concentration of aromatics compared to bio-oil from biomass pyrolysis alone.36 Recently, the combustion behavior of SB pyrolysis bio-oil surrogates mixed with hydrocarbons was investigated, and better combustion behavior was reported.37 These studies demonstrate the potential advantages of using HFO in biomass co-pyrolysis.

The motivation behind this study is to explore SB and HFO co-pyrolysis as a possible bio-oil upgrading method. Different ratios of the two feedstock were used to study their interactions. SB, which is rich in nitrogen and oxygen, is expected to result in a significant amount of nitrogen- and oxygen-containing compounds during pyrolysis. The hydrogen-rich HFO can be a potential co-feed with SB to upgrade the pyrolysis vapors in situ. A tubular furnace pyrolyzer was used to study the effect of co-pyrolysis of SB and HFO on the liquid and solid products. The interactions were studied using a thermogravimetric analyzer (TGA). The effect of changing the geometry of the sample holder on the interactions was also studied in the TGA. The liquid product was then analyzed using a gas chromatography/mass spectrometer (GC/MS). In addition, a kinetic analysis of the process was investigated using the mass loss data obtained from the TGA measurements.

2. Materials and methods

2.1. Preparation and characterization

The SB samples were obtained as described in our previous study.⁷ SB plants were grown from seeds at the KAUST greenhouse for a period of approximately 6 months. The temperature

was maintained at 28 °C during the day and 24 °C at night. The humidity was kept in the range of 50-60% during the growth period to ensure optimal growth conditions. All parts of the plant were utilized in this study. To reduce the high mineral content of the plant and avoid sodium volatilization during pyrolysis, a pretreatment washing method was employed.7 After drying and grinding, SB was placed in deionized water for 8 h at \sim 50 °C in a 1:20 v/v ratio. The biomass was then filtered and dried overnight. The inherent sodium content of SB was 10.28 wt% before treatment and was reduced by almost 55% after the treatment.7 More than 86% of the sodium present in SB was retained in the biochar, thus sodium volatilization at the chosen pyrolysis temperature is not expected to be significant. The particle size of SB used in the experiments was 300 µm. The biochemical composition of SB is provided in Table 1.38 A typical Saudi Arabian heavy fuel oil sample was used, which contained 31 wt% diesel and 69 wt% vacuum residual oil. The SARA (Saturates, Aromatics, Resins, and Asphaltenes) of the HFO used is also available in Table 1.39

Proximate and ultimate analyses of SB and HFO were performed and summarized in Table 1. The proximate analysis was performed following the ASTM E 1131-08 standard method using a TGA Q5000 (TA Instruments, USA).⁴⁰ A sample mass of 5 mg was heated in the presence of N₂ gas with a flowrate of 100 mL min⁻¹. The temperature program employed started from room temperature to 105 °C at a heating rate of 20 °C min⁻¹ and was held constant for five minutes. The temperature was then increased from 105 to 900 °C at 10 °C min⁻¹. Once 900 °C was

 Table 1
 Characterization and composition of SB and HFO on dry basis (data represented as: mean (s.d.))

SB		HFO	
Proximate analysis	(% _{mass})	Proximate analysis	(% _{mass})
Moisture	2.22(0.4)	Moisture	0.00
Volatile matter	84.12 (0.3)	Volatile matter	89.66 (2.7)
Fixed carbon	9.88 (0.6)	Fixed carbon	9.78 (0.3)
Ash	3.78 (0.5)	Ash	1.24 (0.02)
Ultimate analysis	(% _{mass})	Ultimate analysis	(% _{mass})
С	48.53 (1.04)	С	83.21 (2.5)
Н	7.38 (0.2)	Н	10.19 (0.3)
Ν	5.02 (0.2)	Ν	0.12 (0.004)
0	35.29 (1.76)	0	1.37 (0.04)
S	<0.5	S	3.87 (0.1)
Biochemical			
composition ^a	(% _{mass})	SARA ^b	(% _{mass})
Cellulose	46	Saturates	35.71
Hemicellulose	15	Aromatics	40.36
Lignin	2	Resins	13.02
-		Asphaltenes	10.9

^{*a*} Reproduced with permission from Bañuelos *et al.*,³⁸ Renewable Energy; published by Elsevier, 2018. ^{*b*} Reproduced with permission from Alabbad *et al.*,³⁹ Journal of Thermal Analysis and Calorimetry; published by Springer Nature, 2023.

Paper

reached, the gas was switched to O_2 at the same flowrate and held constant at 900 °C for 45 minutes. The elemental analysis was performed in a Thermo Flash 2000 (Thermo Fisher Scientific, U.S.A.) using a sample mass of 2–3 mg. Calibration of the instrument with 2,5-bis(5-*tert*-butyl-2-benzo-oxazol-2-yl) thiophene was performed to quantify the relative amounts of carbon, hydrogen, nitrogen and sulfur in the sample. The oxygen content of the sample was calculated by difference, which is commonly used for the elemental analysis of biomass species. Elemental analyses of biochar samples collected from co-pyrolysis experiments were also performed using the same procedure.

Mixtures of SB and HFO were prepared to study their interactions. The HFO content in the mixtures was 25%, 50% or 75%, with respect to the total dry mass of the sample. Pyrolysis experiments were performed on the individual components as well as the mixtures, resulting in a total of 5 samples: SB or 0% HFO, 25% HFO, 50% HFO, 75% HFO, and 100% HFO.

2.2. TGA experiments

The thermogravimetric analysis of the samples was performed using a TGA Q5000 (TA Instruments, USA). Approximately 5 mg of each sample was used in high-temperature platinum pans. The temperature ranged from ambience to 900 °C at a heating rate of 10 °C min⁻¹ at atmospheric pressure. These experiments were performed under N_2 conditions at a flow rate of 100 mL min⁻¹.

The thermogravimetric analyzer-Fourier transform infrared spectroscopy (TG-FTIR) experiments were performed using a Netzsch TG209 TGA coupled to a Bruker Tensor 27 FTIR. Approximately 5 mg of each sample was placed in open alumina crucibles. The experiments were conducted at atmospheric pressure with the same temperature range and heating rates as in the TGA experiments described above. The evolved vapors were carried to the FTIR *via* a transfer line made of Teflon tubes (800 mm (l) 4 mm (i.d.)), and were maintained at 230 °C to avoid the condensation of the vapors. The detector used was a Mercury–Cadmium–Telluride (MCT) IR detector. The FTIR spectra were collected in the wavenumber range of 650–4500 cm⁻¹ at a scan rate of 2 cm⁻¹ and are provided in the ESI.† Both TGA and TG-FTIR experiments were repeated to establish reproducibility, and negligible variations were observed.

The TGA mass loss profiles from the different sample holders were utilized to study the effect of the free space available above the sample on the mass loss data. A figure of the sample holders along with their dimensions can be found in ESI.[†]

2.3. Non-isothermal kinetic analysis

Thermogravimetric data was used to find the apparent activation energy for the pyrolysis of each sample. The general rate expression for solid-state decomposition is

$$\frac{\mathrm{d}\alpha}{\mathrm{d}t} = A \, \exp\!\left(\frac{-E}{RT}f(\alpha)\right) \tag{1}$$

where *t* is time (min), *A* is the pre-exponential factor (min⁻¹), *E* is the activation energy (J mol⁻¹), *R* is the universal gas constant (J mol⁻¹ K⁻¹), *T* is the temperature (K), and $f(\alpha)$ is the kinetic model function dependent on conversion. The normalized conversion, α , is defined as

$$\alpha = \frac{m_{\rm o} - m}{m_0 - m_{\infty}} \tag{2}$$

where m_0 , m, and m_∞ represent the initial, transient and final mass of sample (mg), respectively. Using the Coats–Redfern integral method, the activation energy was calculated by rearranging and integrating eqn (1), keeping in mind that $\beta = \frac{dT}{dt}$, the heating rate, is constant during the experiment.^{41–43} The following is obtained

$$\ln\left(\frac{g(\alpha)}{T^2}\right) = \ln\left(\frac{AR}{\beta E}\left(1 - \frac{2T}{E}\right)\right) - \frac{E}{RT}$$
(3)

where $g(\alpha)$ is the integral form of the kinetic model function $f(\alpha)$. The kinetic model function is determined from a list of solid state degradation mechanisms (provided in ESI[†]).⁴⁴ The apparent activation energy can be found by plotting $\ln\left(\frac{g(\alpha)}{T^2}\right) vs \frac{1}{T}$ (eqn (3)). $g(\alpha)$ is evaluated using the normalized conversion and corresponding model function and temperature is recorded using a TGA. This will give a straight line with a slope equal to $\frac{-E}{RT}$, from which the apparent activation energy can be evaluated.

2.4. Experimental setup

Pyrolysis experiments were carried out in a customized tubular reactor. The reactor consisted of a quartz tube with stainless steel fittings attached at the ends for the gas inlet and outlet. The tube was placed in an electric furnace CY-O1200-50ICS (CY Scientific Instrument, China). The gas flow rate of N2 was maintained at 300 mL min⁻¹. The reactor was purged with N₂ prior to each experiment to ensure an inert environment. A sample of about 1.8 g was placed in a quartz boat sample holder and pushed into the center of the reactor tube once the desired temperature of 550 °C was reached. The temperature was also monitored by inserting a thermocouple into the reactor. In the mixtures, SB was placed at the bottom of the boat and the corresponding amount of HFO was placed above it, ensuring even distribution along the length of the quartz boat on the SB sample. A two-stage condenser was used at the outlet of the reactor to help condense the vapors and collect the liquid product. A schematic of the reactor setup can be found in ESI.† Each experiment was repeated at least three times nonconsecutively to establish confidence in the data. The actual heating rate experienced by the sample was not determined in this study, however, other studies have shown that a biomass sample in this type of setup typically experiences a heating rate of about 100-200 °C min⁻¹.45 This heating rate estimation will depend on the properties of the sample such as thermal conductivity, specific heat capacity and density. The gaseous product was not collected in this experiment, as the focus of the study was to understand the effect of interactions on the quality of the co-pyrolysis liquid bio-oil.

2.5. Product characterization

The elemental analysis of the solid product (biochar) collected form SB, HFO and their mixtures was performed following the method presented in Section 2.1. The liquid product (bio-oil) collected from the pyrolysis experiments was analyzed in an Agilent 7890A gas chromatograph/mass spectrometer (GC/MS). The organic phase of the bio-oil was diluted with dichloromethane in equal amounts. An Agilent HP-5MS capillary column (60 m length \times 0.25 mm diameter \times 0.25 μ m film thickness) was used for the separation. About 1 µL was injected into the column at an injection temperature of 250 °C. Ultrapure helium (99.9995%) was used to carry the vapors through the column at a flow rate of 2 mL min⁻¹ and a split ratio of 5 : 1. The oven temperature was programmed to increase at a rate of 10 °C min⁻¹ from 35 to 320 °C. The oven was held at 320 °C for 5 min isothermally. The interface and detector temperatures were maintained at 300 °C and 250 °C, respectively. The electron ionization (EI) potential of the ion source was 70 eV and the scanning m/Z range was 35-700 Da. The National Institute of standards and technology (NIST) data library was used to identify the compounds present in the bio-oil based on the mass spectra. To understand the interactions between SB and HFO in the co-pyrolysis experiments, a deviation parameter (D) was introduced and was defined based on the selectivity as

$$D = \left(\frac{S_{\exp} - S_{calc}}{S_{calc}}\right) \times 100 \tag{4}$$

where S_{exp} denotes the experimental selectivity of a specific functional group and S_{calc} denotes the calculated selectivity, which can be found from the following formula

$$S_{\text{calc}} = \frac{m \times S_{\exp(\text{SB})} + n \times S_{\exp(\text{HFO})}}{m + n}$$
(5)

where m and n are the mass ratios of SB and HFO in the mixture.

A positive *D* value represents an increase in selectivity of that group in the mixture ratios under study, which indicates synergistic interactions. A negative *D* value represents a decrease in the selectivity and indicates antagonistic interactions. A similar approach was used by Gautam and Vinu⁴⁶ to study the interactions between binary and ternary mixtures of proteins, lipids and carbohydrates.

3. Results and discussion

3.1. Interactions between SB and HFO during co-pyrolysis in a TGA

3.1.1. TG and DTG profiles. Experimental TG and DTG profiles of the individual components, SB and HFO, are presented in Fig. 1. Both SB and HFO have two general decomposition regions. The first decomposition region in HFO is due to the evaporation of light hydrocarbons (alkanes and alkenes) and can be observed in the DTG profile at temperatures below 200 °C. Heavy hydrocarbons, such as aromatics, resins and asphaltenes, in HFO decompose at higher temperatures between 300-500 °C, as observed in the DTG profile. Similar peaks were observed by Abdul Jameel et al.31 and Alabbad et al.39 for HFO pyrolysis. In biomass, the first decomposition region in the DTG profile is attributed to moisture loss. SB exhibits a wider second decomposition region (150-500 °C) which is due to the complex nature of hemicellulose, cellulose, and lignin. More specifically, hemicellulose and proteins in SB decompose in the temperature range of 200-330 °C. At higher temperatures (300-400 °C), cellulose decomposition occurs. Finally, lignin and lipids decompose in a wider temperature range, 150–900 °C and 290-540 °C, respectively.46 The temperature at which the maximum mass loss rate is observed (T_{gmax}) was 325 °C for SB and 450 °C for HFO. HFO exhibits a higher Tgmax which is attributed to its high content of polycyclic aromatic hydrocarbon pool in the form of resins and asphaltenes.47 The mass remaining beyond 500 °C is higher in SB than HFO due to the higher content of ash in SB, which is also evident in the proximate analysis in Table 1.



Fig. 1 (a) TG and (b) DTG profiles of the individual components pyrolyzed in a high temperature platinum pan at 10 K min⁻¹.



Fig. 2 Theoretical and experimental DTG profiles of the three mixtures pyrolyzed in a high temperature pyrolysis pan (a) 25% HFO, (b) 50% HFO and (c) 75% HFO.

The experimental and theoretical DTG profiles of the three mixtures are shown in Fig. 2. The theoretical values were calculated on a weighted average basis. Theoretical and experimental mass loss profiles overlapped, and no significant differences are observed. The DTG profiles of the mixtures are similar to those of individual SB and HFO, which is consistent with other studies on biomass co-pyrolysis.^{22,48} The $T_{\rm gmax}$ of the individual components, SB and HFO, are observed in the mixtures' DTG profiles as expected. A decrease in the maximum weight loss rate in the first stage and an increase in the second stage is observed when compared to the theoretical values. However, no significant interactions are observed in the DTG profiles. It should be noted that the pyrolysis interactions in a TGA do not capture any heat and mass transfer limitations due to the small sample size and low heating rate. Therefore, it does not provide an accurate representation of the interactions that would take place in a real size reactor.49 This is also supported by literature exploring the co-pyrolysis of cellulose with low density polyethylene and tobacco stalk with polymer.^{22,50}

The pyrolysis process in a TGA is kinetically controlled and the TGA data collected was used for apparent kinetic analysis to understand the reactivity of SB and HFO mixtures, following the ICTAC Kinetics Committee recommendations.⁵¹ To further investigate the effect of mixing, other analyses such as TG-FTIR and GC/MS were also performed.

A set of co-pyrolysis TG experiments were performed in an alumina crucible and a platinum pan (see ESI† for details). Studies have shown that the size and shape of the sample holder and the free space volume available can affect the mass loss profiles in TGA.^{52,53} To have uniformity in the measurements, the sample mass used in these experiments was 5 ± 0.5 mg. The two sample holders differ in the free space volume available above the sample. The alumina crucible offered larger free space volume compared to the pan. Fig. 3 shows the DTG curves obtained from the decomposition of SB and HFO in the alumina crucible and the platinum pan.

The onset decomposition temperature of HFO shifted to higher temperatures using the alumina crucible. The



Fig. 3 DTG curves of SB and HFO in alumina crucibles and platinum pans.

temperature shifts from around 250 °C in the pan to 350 °C in the crucible. The peak temperature, however, remains the same. This could be explained by a delay in carrying out the vapors formed on top of the sample due to the difference in the height of the sample holder. In the pan, the purging gas can easily carry out the vapors, unlike the alumina crucible. Similar results have been observed by Shi *et al.* in which they have designed experiments to study the effect of the geometry of the crucible on thermal analysis.⁵⁴ In SB the onset decomposition temperature has also slightly shifted to higher temperatures in the crucible. This is also due to delays in carrying out the pyrolysis vapors.

The peak temperature height was also affected by changing the sample holder. SB showed a higher peak height in the pan. This again could be due to the height of the walls. The free space available increases the diffusion pathway of the gas and hence results in a lower apparent rate. This was also observed by Nowak *et al.*, as the free space available was altered by the addition of Al_2O_3 below the biomass sample.⁵⁵ A lower rate was observed when the diffusion pathway through the free space was higher. In contrast, the opposite was observed in HFO. For HFO, a higher peak height was observed in the alumina crucible. This observation could be related to the concentration and rate of reaction. The initial concentration of the material at the decomposition temperature is lower in the pan compared to the crucible due to easier flashing of the lighter cut portion of HFO resulting in a lower apparent rate in the pan.

The DTG of the mixtures follow the characteristics observed in SB and HFO and any differences observed follow the same reasoning mentioned above. The DTG of mixtures can be found in ESI.[†]

The maximum weight loss of SB in the DTG profile was around 325 °C in both crucibles (Fig. 3). The evolution of gases around the peak temperature corresponds to cellulose, hemicellulose and protein decomposition.⁴⁶ Around this temperature, depolymerization, cleavage of the glycosidic bond fragmentation, dehydration and rearrangement reactions occur. These reactions result in compounds such as furfural and benzene derivatives which evolve from cellulose decomposition. Whereas, hemicellulose decomposition results in furan derivatives, acetaldehyde and hydroxyacetaldehyde. Finally, protein decomposition results in compounds containing pyridine and indole as well as aldehydes and benzamine. Some of these compounds and reactions are represented in Fig. 7b. These reactions reduced as the composition of HFO increased in the mixture. Further discussion can be found in the TG-FTIR section in the light of product formation.

3.1.2. TG - FTIR. FTIR spectra in the temperature range of 105-600 °C are presented in ESI.[†] From the FTIR spectra, the deconstruction of biomolecules present in the biomass started around 200 °C. The onset of CO2 evolution from biomass pyrolysis has been reported after 200 °C, and the time for maximum vapor evolution for the majority of the functional groups was observed between 300-400 °C in a TGA-FTIR.56 Protein, cellulose and hemicellulose have been reported to undergo thermal decomposition reactions to produce CO₂, carbonyls and other oxygenates in this temperature range. The early evolution of C-H stretch in the TGA-FTIR spectra of HFO can be attributed to the release of lighter hydrocarbons present. Characteristic peaks representing the presence of various aromatic compounds were also seen in the FTIR spectra, which is consistent with the literature.³¹ However, the main functional groups observed are discussed in this section.

Fig. 4 depicts the major functional groups observed in the pyrolysis vapors of SB, HFO, and their mixtures. The major groups include CO_2 (2350 cm⁻¹), methyl C–H stretch (2935 cm⁻¹), aromatic C–H in-plane-bend (906 cm⁻¹) and aromatic C–H out-of-plane bend (690 cm⁻¹). The maximum absorbance peak for CO_2 was observed around 350 °C for all samples except 100% HFO. The intensity of the CO_2 peak was highest for SB pyrolysis. A similar CO_2 peak was observed upon pyrolysis of Tobacco stalk with a maximum at 350 °C.⁵⁰ CO_2 is a major gas produced during biomass pyrolysis due to the presence of oxygenated biomolecules, and it is also formed



Fig. 4 Temperature evolution of major function groups in the pyrolysis products from SB, HFO and their mixtures

from the decarboxylation of carboxylic acids.⁵⁷ The intensity of the CO₂ peak decreased as more HFO was added to the mixture, until no CO₂ was observed in the pyrolysis vapors of 100% HFO. This is expected, as HFO is primarily composed of carbon and hydrogen and lacks oxygen, which is required for the formation of CO₂. It can be noted that CO₂ peaked in the temperature range 340–345 °C for SB and SB/HFO mixtures. As the HFO composition increased, the experimental peak heights were 9.75% (25% HFO), 27.8% (50% HFO) and 46.67% (75% HFO) higher compared to theoretical peak heights indicating promotion of decarboxylation reactions under co-pyrolysis conditions.

HFO decomposes into different aliphatic and aromatic hydrocarbons under an inert environment,³¹ which is evident from the presence of peaks around 2935 cm⁻¹, 906 cm⁻¹ and 690 cm⁻¹ in the FTIR spectra. These peaks correspond to the vibrations of aliphatic and aromatic hydrocarbons, respectively. Aliphatic hydrocarbons were produced from the pyrolysis of all five samples, with the highest intensity peak observed for HFO followed by the 75% HFO mixture. The gas generation of these

hydrocarbons was concentrated between 400 and 650 °C for all samples except SB, which had a wider temperature range of 200–650 °C. This wider temperature range corresponds to the slow degradation of the biomass, as discussed earlier. Mixing of SB with HFO showed no effect on the evolution of aliphatic hydrocarbons as seen in Fig. 4. This suggests that the mixing of SB and HFO does not significantly alter the production of aliphatic hydrocarbons during co-pyrolysis.

Aromatic C–H in-plane bending, which is associated with the presence of polycyclic aromatic hydrocarbons (PAH) such as naphthalene,⁵⁸ was present in the vapors produced from the pyrolysis of SB, HFO and their mixtures. PAH can form from phenol derivatives in lignin when hydroxyl and methoxy linkages are cleaved.⁵⁹ The low intensity of this peak in SB can be attributed to the low content of lignin, which is around 2% by weight. HFO contains non-protonated aromatic carbons, which indicate the presence of fused rings.³¹ The mixtures exhibited a higher intensity peak than that from HFO alone, suggesting that the addition of SB helped in the extraction of the PAH present in HFO. This confirms the synergistic effects of mixing

SB with HFO. The 75% HFO mixture resulted in the highest intensity peak, possibly due to the optimal balance of lignin and HFO content.

C–H out of plane bending, which is associated with the presence of monocyclic aromatic hydrocarbons (MAH), was not observed in HFO pyrolysis. The absence of MAH indicates the presence of more complex aromatics, such as polycyclic aromatic hydrocarbons (PAH). The intensity of the peak is highest for SB, which contains a higher percentage of MAH than HFO, and gradually decreases as the percentage of HFO added increases in the mixture until none is observed at 100% HFO.

3.2. Kinetic analysis

Eqn (3) was used to calculate the apparent activation energy (E_{α}) of the pyrolysis process for each sample. The apparent activation energy can be determined from the slope of the lines produced from plotting the logarithmic term, $\ln\left(\frac{g(\alpha)}{T^2}\right)$, against $\frac{1}{T}$. The plot obtained from eqn (3) was split into three pyrolysis stages based on linearity. Multiple nucleation-growth kinetic models were tested in each stage to find the best fit model. The first stage of each sample was well represented by a model of the form, $g(\alpha) = -\ln(1 - \alpha)$, which describes the initial decomposition of the sample. The second and third stages were both sufficiently represented by a model of the form $g(\alpha) = [-\ln(1 - \alpha)]^4$, which describes the later stages of the decomposition process. Table 2 summarizes the E_{α} values for all samples calculated using these models.

In TGA, the main objective is to perform measurements in a kinetically controlled regime. This is achieved using a low heating rate, small sample mass, and small particle size to ensure that the pyrolysis is kinetically controlled. The low heating rates also help to ensure that there is no (or minimal) gradient in the sample temperature and furnace temperature. In this study, the particle size of SB was 300 μ m, sample mass was around 5 mg and heating rate was 10 °C min⁻¹. The kinetic regime was verified by changing the sample mass and acquiring

 Table 2
 Summary of the activation energy of individual components and their mixtures for each stage

	Stage	Temperature (°C)	Conversion (α)	$E_{\rm a}$ (kJ mol ⁻¹
0% HFO	1	25-62	0-0.03	39.18
0701110	2	62-227	0.03-0.14	35.98
	3	227-500	0.14-0.89	126.96
25% HFO	1	25-74	0-0.03	33.34
	2	74-234	0.03-0.18	59.62
	3	234-500	0.18-0.93	125.03
50% HFO	1	25-106	0-0.06	33.56
	2	106-286	0.06-0.28	52.72
	3	286-500	0.28-0.85	117.31
75% HFO	1	25-123	0-0.09	36.59
	2	123-358	0.09 - 0.48	55.01
	3	358-500	0.48-0.96	191.62
100% HFO	1	25-73	0-0.05	51.15
	2	73-356	0.05 - 0.46	43.10
	3	356-500	0.46 - 0.97	200.26

mass loss profiles. These overlapping mass loss profiles established the kinetically controlled regime for the performed TGA experiments. A detailed discussion can be found in literature.^{60–62} The main objective of investigating the nonisothermal kinetics in this study is to understand the shifts in different temperature regimes observed with mixing.

The thermal decomposition of the biomass, HFO and their mixtures followed a sigmoidal reaction regime. The kinetics of these reactions were well-captured by the nucleation-growth model. The kinetic functions used to evaluate the apparent activation energy represent both random nucleation and sequential growth mechanism. The corresponding equation leads to a sigmoidal reaction progression, as observed in many solid-state reactions, including this study. Solid-state reactions during pyrolysis commonly occur by the generation of nucleation sites followed by their geometric growth. It is worth noting that nucleation-growth models for solid-state reactions have been widely used in the last two decades to model the pyrolysis of organic materials.⁶³

Decomposition occurs at higher temperatures, and therefore it is expected to observe higher E_{α} at higher temperatures. This is confirmed by the higher E_{α} values observed at stage 3 of all five samples compared to stages 1 and 2. HFO exhibited a higher E_{α} value compared to SB in all three stages. This can be attributed to its high thermal stability, which is due to the presence of asphaltenes that require more energy to decompose.

Biomass degradation takes longer, as most of the decomposition occurs in stage 3, where the conversion jumps from 0.14 to 0.82. However, HFO reaches almost 50% conversion in the second stage. The effect of this observation can also be extended to the mixtures, where higher percentages of HFO result in higher conversions at earlier stages corresponding to lower temperatures.

In stage 1, mixing resulted in slightly lower E_{α} values compared to the individual components. In stage 2, the mixtures showed slightly higher E_{α} values than SB and HFO individually, but the effect of mixing was not significant in these stages. The first stage of the mass loss corresponds to the release of moisture from the biomass and lighter components from the HFO, which is characterized by low E_{α} values. In stage 2, the decomposition of cellulose, hemicellulose and proteins begins, which is consistent with previous.46 In stage 3, the mixtures exhibited lower E_{α} values the individual components. Interestingly, E_{α} of HFO can be significantly reduced by about 42% by equal mixing with SB. This higher reactivity in the third stage could be attributed to biomass radicals potentially initiating reactions and enhancing the decomposition of HFO leading to lower E_{α} values. This finding is supported by previous studies that suggest biomass can act as a radical initiator in pyrolysis reactions.48

3.3. Interactions between SB and HFO during co-pyrolysis in tubular reactor

3.3.1. Product yields. The product yields from each experiment are summarized in Fig. 5. The highest char yield, around

Paper

21%, was obtained from the pyrolysis of SB alone. The yield of solid decreased gradually as the percentage of HFO increased, with the lowest char yield of 13% obtained at 100% HFO. This observation can be attributed to the ash content in the individual components. Since SB has higher ash and fixed carbon content, it exhibits a higher char yield after pyrolysis. In contrast, the liquid yield showed the opposite trend, gradually increasing from 52.6% to 67.4% as the percentage of HFO increased from 0% to 100% in the samples. This increase can be linked to the slightly higher amount of volatile matter in HFO. It should be noted that the fuel properties of the bio-oil were not investigated in this study due to the nature of the experiment. Considering the size of the bench-scale reactor, the samples used in the experiments were small (around 1.8 g) and the biooil collected was recovered by diluting it with DCM as it tends to sticks to the walls of the collection apparatus.

To study the interactions during co-pyrolysis, the experimental yields were compared to the theoretical values calculated on a weighted average basis of the individual components. Synergistic interactions were observed in the gas and char yields. Mixing resulted in increased production of gas compared to the theoretical value, indicating a synergistic interaction. Similarly, the experimental char yields were lower than the theoretical values, which indicates that mixing enhanced devolatilization of the sample and allowed for a higher degree of conversion of the original feedstock. A hydrogen rich environment provided by HFO creates more radicals by transferring hydrogen to the biomass. These radicals initiate and promote scission and cracking reactions, which enhance char decomposition and increase gas yields. This phenomenon has been observed in previous studies.^{24,48,64}

3.3.2. Elemental analysis of solid residue. The elemental analyses of the solid residue collected from the five co-pyrolysis experiments are presented in Table 3. Carbon was enriched in SB biochar and HFO solid residue by about 12% and 3%, respectively, compared to the carbon content of the feedstock. The carbon and sulfur content in the solid residue increased as the percentage of HFO added increased, whereas the other elements (H, N, and O) decreased.



Fig. 5 (a) Char, (b) gas and (c) liquid product yields of the pyrolysis of SB, HFO and their mixtures.

Table 3	Elemental analysis of solid resi	due from SB, HFO an	d the mixtures on dry ash	n free basis (data represented	as: mean (s.d.))
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Sample	С	Н	Ν	S	0
SB	61.23 (2.21)	3.83 (0.02)	6.98 (0.54)	n.d.	27.96 (2.58)
25% HFO	66.78 (3.89)	3.31 (0.01)	5.52 (0.12)	2.28 (0.13)	22.11 (1.15)
50% HFO	75.15 (3.03)	2.68 (0.05)	3.82 (0.22)	4.58 (0.15)	13.77 (3.45)
75% HFO	79.52 (0.60)	2.08 (0.17)	2.48 (0.08)	6.96 (0.87)	8.96 (0.52)
100% HFO	86.94 (2.4)	1.62 (0.04)	0.92 (0.07)	9.38 (0.09)	1.14 (2.47)

Carbon enrichment occurs *via* condensation and aromatization reactions. That involve the decarboxylation, dehydration, and dehydrogenation of the precursors.⁶⁵ These reactions are accompanied by the reduction of nitrogen-containing compounds.⁶⁶ The observed trend of increased carbon content in the solid residue accompanied by a decrease in H, N and O indicates that the addition of HFO might have enhanced these reactions.

3.3.3. GC/MS of liquids. Over 100 compounds were identified for each of the five liquid samples collected. Compounds were identified against a standard mass spectral library, and those with a match factor above 85% were considered for analysis. They were broadly classified based on their functional groups into the following 9 categories: aliphatic hydrocarbons, aromatic hydrocarbons, carboxylic acids, N-containing compounds, esters, carbonyls (aldehydes and ketones), alcohols, phenols, and furans. A detailed list can be found in the ESI.[†]

Fig. 6 represents the selectivities of the major functional groups identified in the bio-oil from pyrolysis of SB, HFO and their mixtures along with the deviation in selectivities calculated using eqn (4). The compounds in the bio-oil can be linked to the components of SB and HFO.

SB has typical characteristics of lignocellulosic biomass, which is ~46 wt% cellulose, ~15 wt% hemicellulose ~2 wt% lignin.³⁸ Lipids and protiens are also present in the biomass matrix. The major amino acids present are proline, phenylalanin, leucine, glutamic acid, tyrosine, histidine and aspartic acid.^{67,68} Lipids are represented by triglycerides in SB. HFO, on the other hand, is a mixture of polycyclic aromatic hydrocarbons containing sulfur, oxygen and nitrogen as heteroatoms. A surrogate has been used to represent HFO containing compounds such as naphthalene, benzothiophene, dibenzothiophene and quinoline.⁴⁷ Fig. 7 shows a representative composition of SB and HFO, as well as a broad overview of the reactions under pyrolysis and co-pyrolysis conditions.

The addition of HFO resulted in a general increase in peak area for aliphatic and aromatic hydrocarbons in the mixture. The peak area % of aliphatic and aromatic hydrocarbons increased by 38% and 21%, respectively, when 75% HFO was added to SB. This increase was accompanied by a positive deviation in the selectivities for these two categories, indicating synergistic interactions. The positive deviation can be attributed to enhanced extraction of fatty acids assisted by pyrolysis and their subsequent decarboxylation in the presence of HFO, resulting in an increase in aliphatic hydrocarbons. Aromatic hydrocarbons, on the other hand, come from lignin dehydration, decarboxylation and demethylation reactions that are also enhanced by the addition of HFO. Furthermore, HFO cracking also contributes to the amount of aromatic hydrocarbons present in the bio-oil. The presence of HFO aids in the generation of more free radicals during co-pyrolysis, which facilitates cracking of C–C bonds and enhances the quality of oil products. The major hydrocarbons detected were eicosane and cetene, as well as benzene and naphthalene derivatives.

In contrast, carboxylic acids exhibit a decrease in peak area, and a negative deviation parameter was recorded as the percentage of HFO added increased. The area % of carboxylic acids from SB pyrolysis was 36%, which dropped to 2% in the 75% HFO mixture, with a deviation of about -81.2%. Cracking of lipids during the pyrolysis of SB results in the generation of fatty acids which are then consumed through decarboxylation reactions resulting in the formation of hydrocarbons and carbon dioxide. The decrease in area% of carboxylic acids confirmed that decarboxylation reactions were enhanced with the addition of HFO. The major acids found in the co-pyrolysis bio-oil were palmitic acid, stearic acid, oleic acid and linoleic acid.

N-containing groups exhibited a negative deviation parameter. This category includs amines/amides, N-aromatic compounds and nitriles. N-containing compounds come from protein cracking, as proteins are the major source of nitorgen in the mixture.46,69 Possible amino acid reaction pathways to produce a variety of N-conatining compounds are presented in Fig. 7b.70 For example, the benzene ring in tyrosine and phenylalanine and the five member ring with nitrogen in proline can undergo decarboxylation and dehydration reactions to form some of the major N-containing compounds present in the bio-oil, such as Indole and Imidazole. The major decomposition pathways of common amino acids in the protein structure of biomass during pyrolysis have been studied and correspond to those found in SB pyrolysis.71 Strecker degradation reactions of some proteins can form the aldehydes found in the bio-oil as seen in Fig. 7b. SB is known to be a protein-rich biomass as seen from the high N-content, which corresponded to a high area% of N-containing compounds in the bio-oil from SB.7,8,67,68 Lack of N-containing compounds in the 100% HFO sample asserts that only SB pyrolysis results in these species. Therefore, a decrease in the composition of this group is expected as more HFO is added. Mixing also contributed to the decrease, as evident in the negative deviation parameter. For example, the 25% HFO sample exhibited a deviation parameter of around -49.3%. The addition of HFO inhibited dehydration and decarboxylation of amino acids that would have allowed for



Fig. 6 Distribution of compounds in the bio-oil from pyrolysis of SB, HFO and their mixtures.

the formation of N-containing compounds from the proteins present in the biomass. Examples of the compounds found in the bio-oil are amides such as oleamide and aromatics such as indole and 2,5-dimethylpyridine. Nitiriles were also present, such as hexadecanitrile, octadecanitrile and benzenepropanenitrile. Amides and nitriles were formed due to the reactions between the pyrolysis products from proteins and lipids.^{8,68}

Other catagories represented only 1–3 area%, and therfore, were not included in the product analysis. However, a detailed list of products is provided in the SM. These catagories include esters such as methyl palmitate and oxalic acid (2-ehtyl hexyl) ester, alcohols such as palmityl and linoleyl alcohol, phenols such as p-cresol and finally furans such as 2-vinylfuran and 2,5furandione. All of which were present in the bio-oil in trace amounts.

3.3.4. Reaction scheme. The reactions involved in the copyrolysis of SB and HFO are difficult to track due to the complexity of the two components and the wide variety of compounds that result from their co-pyrolysis. Fig. 7a shows the molecular composition of biomass, namely, cellulose, hemicellulose, lignin, lipids and proteins as well as the composition of HFO. Lipids are represented by triglycerides and the major amino acids in SB are used to represent proteins.^{67,68,72} The detailed list of the amino acids and fatty acids present in SB can be found in ESI.[†] A surrogate was used to represent HFO.⁴⁷ The lighter cut components present in HFO were also represented in



Fig. 7 (a) Composition of SB and HFO. (b) Plausible reaction scheme for co-pyrolysis of SB and HFO.

the composition of HFO.⁷³ These compounds include normal/ branching alkanes, single ring cycloalkanes and 2 ring aromatics. The lighter cut components of HFO start to vaporize at lower temperatures than those required for pyrolysis. Therefore, some hydrocarbons flash directly and are present in the liquid collected. This was also observed during the pyrolysis

Paper

of HFO at 400 and 500 °C.74 The main reactions observed during HFO pyrolysis were cracking of non-condensed rings leading to the significant formation of monocyclic aromatics and thiophene derivatives. Also, reactions such as homolytic bond fission and hydrogen abstraction creating radicals that then undergo B-scission producing unsaturated hydrocarbons or radical addition to generate long chain radicals and finally radical termination via recombination creating longer chains occur.39 These reactions result in compounds such as aliphatic hydrocarbons, naphthalene and benzothiophene derivatives. Fig. 7b provides a broad overview of the possible reactions involved in the co-pyrolysis process based on the major constituents of biomass and HFO. These reaction pathways were predicted based on the known structure and composition of SB and HFO as well as the compounds present in the bio-oil after pyrolysis.

The primary pathways for biomass decomposition are cracking, depolymerization, rearrangement and char formation, with secondary reactions involving cracking and recombination also contributing to the process.⁷⁵ Based on the GC/MS analysis of the bio-oil presented in the previous section, blue arrows indicate the reactions that were enhanced by the addition of HFO, while red arrows represent those that were inhibited.

During co-pyrolysis, the hydrocarbons generated from the cracking of HFO undergo chain-end and random scission, leading to the transfer of hydrogen to the anhydrosugars produced from cellulose pyrolysis and the formation carbonyl compounds such as aldehydes and ketones. The co-pyrolysis process promotes the conversion of aldehydes and ketones to hydrocarbons. Oxygenated compounds produced from cellulose pyrolysis act as acceptors, facilitating the scission of hydrocarbons into smaller alkanes and alkenes. As shown in Fig. 7, an increase in the biomass composition in the feedstock promotes the production of aromatic hydrocarbons via Diels-Alder reaction.⁷⁶ Furan derivatives and other light oxygenates produced from biomass pyrolysis are consumed by the olefins produced from HFO pyrolysis. It is important to note that the interactions between the vapors from both the feedstocks during co-pyrolysis is known to be a complex phenomenon. If the biomass composition in the feedstock is too low or too high, it can significantly hinder the interactions between biomass and co-feed. This is why moderate compositions of HFO and biomass were chosen for this study.

A thorough analysis of gas and liquid samples would provide further insights into the interactions between biomass and HFO during co-pyrolysis, and this can be achieved by increasing the sample mass. The information from this study can be used, in part, for investigation of gasification of HFO with biomass as cofeed. Blends of pyrolysis bio-oil with heavy fuel oil have been shown to have good stability, reduced viscosity, and inhibition of polymerization.⁷⁷ Although direct information on the properties of bio-oil obtained from pyrolysis of biomass with HFO is not available, studies elsewhere have compared the properties of bio-oil from co-pyrolysis of biomass with a hydrogen rich source (*e.g.*, plastics) to that of bio-oil from biomass pyrolysis. For instance, bio-oil produced from the co-pyrolysis of rice straw with polyethylene waste exhibited lower viscosities and higher calorific values compared to bio-oil produced from biomass pyrolysis. The flash points of the bio-oil from mixtures were slightly higher than that of HFO and were comparable to that of diesel.⁷⁸ Based on this, some enhancement in the fuel properties of the bio-oil from co-pyrolysis of SB and HFO is expected, suggesting their potential utilization for reduced emissions.

4. Conclusion

The effects of mixing SB biomass with HFO during pyrolysis at 550 °C were understood through experiments performed in a TGA and a tubular reactor. The free space available in the two sample holders used had some effect on the mass loss data obtained from the TGA. This was mainly due to the obstruction from the walls of the alumina crucibles in carrying out the vapors produced during pyrolysis. The results of the co-pyrolysis experiments show some synergistic effects between the two components. The addition of HFO resulted in increased extraction of PAH in the vapors produced during the pyrolysis. A kinetic analysis of the process revealed that the addition of HFO lowered the apparent activation energy in stage 3 (227-500 °C) of the co-pyrolysis process. Mixing also promoted cracking and volatilization hence increased gas yields by 7% and decreased char yields by 13% compared to the theoretical value of the 75% HFO mixture. A GC/MS analysis of the collected bio-oil showed an increase in selectivity towards aliphatic and aromatic hydrocarbons and a decrease towards carboxylic acids. The peak area% of aliphatic and aromatic hydrocarbons increased by 38% and 21%, respectively, and decreased by 34% for carboxylic acids when 75% HFO was added to the mixture. This study shows the benefits of co-pyrolysis of SB and HFO to produce valuable chemicals and possible upgrading of the fuel.

Conflicts of interest

Authors have no competing financial interests to declare.

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