Entrained flow reactor (EFR)

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1 Introduction

This report provides an overview of the Entrained Flow Reactor (EFR) at NREL and associated computational modeling tasks. The reactor operates at fast pyrolysis conditions to thermochemically convert biomass into gaseous products. The EFR is part of the Thermochemical Process Development Unit (TCPDU) at NREL which was originally designed for biomass gasification where the EFR was used as a thermal cracker. An overview of the TCPDU system is shown in Figure 1.

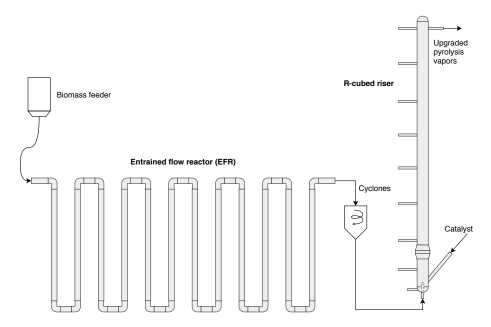


Figure 1: Overview of the main components of the NREL TCPDU system. Fast pyrolysis of biomass occurs in the entrained flow reactor. Catalytic vapor phase upgrading occurs in the R-cubed riser reactor.

2 Experimental setup

This section provides geometric dimensions and typical operating conditions for the entrained flow reactor. Characteristics for the Blend3 and forest residue feedstocks are also discussed.

2.1 Entrained flow reactor

Fast pyrolysis in the TCPDU system occurs in the entrained flow reactor (EFR) which is comprised of a series of horizontal and vertical pipes connected with 90 degree elbows (see Figure 2). The EFR is essentially a pneumatic conveyor

where biomass particles flow through a long pipe with several bends. Dimensions and material information about the EFR are provided in Figure 3 below. Operating conditions such as temperatures, pressures, and flow rates for the EFR are shown in Figure 4. Nitrogen gas at 500°C is generally used as the conveying medium for the solids.

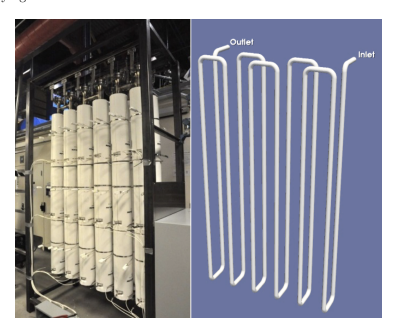


Figure 2: Left - picture of the EFR assembly with heat jackets, insulation, and thermocouples. Right - CAD representation of the EFR pipe assembly used for MFiX simulations.

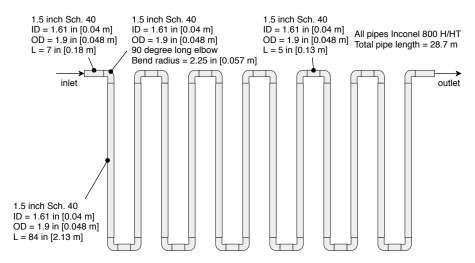


Figure 3: Geometry of the entrained flow reactor at NREL.

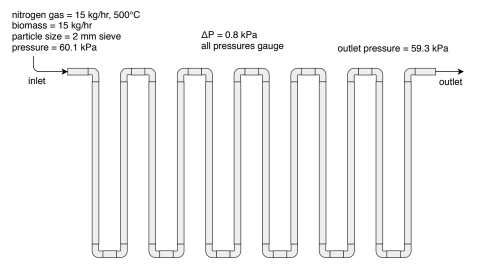


Figure 4: Typical operating conditions for the entrained flow reactor.

2.2 Blend3 feedstock

General information about the Blend3 feedstock used in the entrained flow reactor is provided in Table 1. There is currently no information regarding identification of the feedstock or who performed the feedstock measurements and data preparation. Proximate and ultimate analysis data for the feedstock are presented in Tables 2 and 3. Only one set of analysis data is available therefore the uncertainty in the values is unknown.

Table 1: General information for the Blend3 feedstock.

| Item | Description |
|---------|-------------|
| Name | Blend3 |
| ID | ? |
| Contact | ? |

Table 2: Blend3 proximate analysis mass percent, as-received basis. Source [3].

| Proximate | %ar | %ar | %ar |
|-----------|-------|-----|-----|
| FC | 16.92 | ? | ? |
| VM | 76.40 | ? | ? |
| ash | 0.64 | ? | ? |
| moisture | 6.04 | ? | ? |

Table 3: Blend3 ultimate analysis mass percent, as-received basis. Source [3].

| Element | %ar | %ar | % ar |
|----------|-------|-----|------|
| С | 49.52 | ? | ? |
| ${ m H}$ | 5.28 | ? | ? |
| O | 38.35 | ? | ? |
| N | 0.15 | ? | ? |
| S | 0.02 | ? | ? |
| ash | 0.64 | ? | ? |
| moisture | 6.04 | ? | ? |

The chemical analysis of the Blend3 feedstock is presented in Table 4. Again, only one set of data is available so the uncertainty in the measurements is unknown. The chemical analysis measurements are used to determine the biomass composition which is needed for the kinetics model.

Table 4: Blend3 chemical analysis mass percent, dry basis. Source [8].

| Chemical component | % dry | % dry | % dry |
|---------------------------|-------|-------|-------|
| glucan | 38.95 | ? | ? |
| acetyl | 1.59 | ? | ? |
| arabinan | 1.40 | ? | ? |
| galactan | 3.16 | ? | ? |
| mannan | 10.52 | ? | ? |
| xylan | 7.89 | ? | ? |
| lignin | 29.48 | ? | ? |
| free fructose | 0.07 | ? | ? |
| free glucose | 0.04 | ? | ? |
| sucrose | 0.04 | ? | ? |
| water extractives | 2.75 | ? | ? |
| ethanol extractives | 3.49 | ? | ? |
| non-structural inorganics | 0.22 | ? | ? |
| structural inorganics | 0.41 | ? | ? |

Table 5: Blend3 ash analysis as weight percent of ash. Source [3].

| Metal oxide | wt. % | wt. % | wt. % |
|--------------|-------|-------|-------|
| SiO_2 | 28.1 | ? | ? |
| Al_2O_3 | 7.06 | ? | ? |
| ${ m TiO_2}$ | 0.34 | ? | ? |
| CaO | 21.8 | ? | ? |
| Na_2O | 0.71 | ? | ? |
| K_2O | 13.8 | ? | ? |
| P_2O_5 | 5.47 | ? | ? |
| SO_3 | 1.23 | ? | ? |
| Cl | 0.09 | ? | ? |
| CO_2 | 5.14 | ? | ? |

Table 6: Blend3 particle properties from pelletized crushed feedstock. The crushed feedstock is used in the entrained flow reactor.

| Property | Value | Description | Source |
|------------|--|---|--------|
| ρ | | particle density, daf basis | [7] |
| $\eta \ k$ | $\begin{array}{c} 0.27 \\ 0.23 \; \mathrm{W/mK} \end{array}$ | particle porosity thermal conductivity | |

Table 7: Entrained flow reactor yields for Blend3 feedstock.

| Yield | wt. % |
|----------------|----------------|
| total liquid | 64.9 |
| char | 13.9 ± 0.1 |
| gas | 17.2 ± 0.2 |
| mass balance | 96.9 ± 1.5 |
| carbon balance | 93.0 ± 1.0 |

2.3 Forest residue feedstock

The forest residue feedstock is comprised of branches/twigs, cambium, needles, bark, and whitewood. This feedstock is used in the NREL fluidized bed reactor (FBR) for the purposes of the FCIC project. The FBR is operated at fast pyrolysis conditions for the thermochemical conversion of biomass. The reactor is sometimes referred to as the 2FBR.

Table 8: General information for the forest residue feedstock.

| Item | Description | |
|-----------------------|--------------------|--|
| Name ID Contact | forest residue ? ? | |

Table 9: Bark ultimate analysis mass percent, dry ash-free basis. Source [1].

| Element | %daf | %daf | %daf |
|---------|-------|------|------|
| С | 48.27 | ? | ? |
| H | 5.72 | ? | ? |
| N | 0.52 | ? | ? |

Table 10: Branches/twigs ultimate analysis mass percent, dry ash-free basis. Source [1].

| Element | % daf | % daf | % daf |
|---------|-------|-------|-------|
| С | 49.69 | ? | ? |
| Н | 6.36 | ? | ? |
| N | 0.25 | ? | ? |

Table 11: Cambium ultimate analysis mass percent, dry ash-free basis. Source [1].

| Element | % daf | % daf | % daf |
|---------|-------|-------|-------|
| С | 48.52 | ? | ? |
| Η | 6.39 | ? | ? |
| N | 0.11 | ? | ? |

Table 12: Needles ultimate analysis mass percent, dry ash-free basis. Source [1].

| Element | % daf | % daf | % daf |
|--------------|-------|-------|-------|
| \mathbf{C} | 48.59 | ? | ? |
| ${ m H}$ | 5.92 | ? | ? |
| N | 1.22 | ? | ? |

Table 13: Whitewood ultimate analysis mass percent, dry ash-free basis. Source [1].

| Element | % daf | % daf | % daf |
|--------------|-------|-------|-------|
| С | 48.27 | ? | ? |
| \mathbf{H} | 6.15 | ? | ? |
| N | 0.10 | ? | ? |

Table 14: Whitewood biomass composition mass percent, dry basis. Source [2].

| Component | % dry |
|---------------|-------|
| Cellulose | 38.04 |
| Hemicellulose | 24.2 |

3 Model development

Details about the biomass pyrolysis kinetics, biomass characterization method, and computational models developed for the entrained flow reactor are discussed in the following sections.

3.1 Pyrolysis kinetics

The kinetic reaction mechanisms presented in the Debiagi et al. 2018 paper are used to model biomass pyrolysis in the entrained flow reactor [4]. Table 15

summarizes the reactions along with the associated prefactors and activation energies. A description of the chemical species in the Debiagi et al. kinetic scheme is provided in Table 16. Species are grouped into solid, metaplastic, gas, and liquid phases.

Table 15: Kinetic reactions for biomass pyrolysis where A is the prefactor, E is the activation energy, and T is temperature. Source [4].

| Item | Reaction | A (1/s) | E (cal/mol) |
|------|---|-------------------------------|-------------|
| 1 | $CELL \rightarrow CELLA$ | 1.5×10^{14} | 47,000 |
| 2 | CELLA \rightarrow 0.40 CH2OHCHO + 0.03 CHOCHO + 0.17 CH3CHO + 0.25 | 2.5×10^{6} | 19,100 |
| | C6H6O3 + 0.35 C2H5CHO + 0.20 CH3OH + 0.15 CH2O + 0.49 CO + 0.05 | | |
| | $G{CO} + 0.43 CO2 + 0.13 H2 + 0.93 H2O + 0.05 G{COH2} loose + 0.02$ | | |
| | $\text{HCOOH} + 0.05 \text{ CH2OHCH2CHO} + 0.05 \text{ CH4} + 0.1 \text{ G}\{\text{H2}\} + 0.66 \text{ CHAR}$ | | |
| 3 | $CELLA \rightarrow C6H10O5$ | $3.3 \times T_{2}$ | 10,000 |
| 4 | CELL \rightarrow 4.45 H2O + 5.45 CHAR + 0.12 G{COH2} stiff + 0.18 G{COH2} | 9.0×10^{7} | 31,000 |
| | loose $+ 0.25 \text{ G}\{\text{CO}\} + 0.125 \text{ G}\{\text{H2}\} + 0.125 \text{ H2}$ | | |
| 5 | $GMSW \rightarrow 0.70 \text{ HCE1} + 0.30 \text{ HCE2}$ | 1.0×10^{10} | 31,000 |
| 6 | $XYHW \rightarrow 0.35 \text{ HCE1} + 0.65 \text{ HCE2}$ | 1.25×10^{11} | 31,400 |
| 7 | $XYGR \rightarrow 0.12 \text{ HCE1} + 0.88 \text{ HCE2}$ | 1.25×10^{11} | 30,000 |
| 8 | $\text{HCE1} \rightarrow 0.25 \text{ C5H8O4} + 0.25 \text{ C6H10O5} + 0.16 \text{ FURFURAL} + 0.13 \text{ C6H6O3}$ | $16.0 \times T$ | 12,900 |
| | + 0.09 CO2 + 0.1 CH4 + 0.54 H2O + 0.06 CH2OHCH2CHO + 0.1 CHOCHO | | |
| 0 | + 0.02 H2 + 0.1 CHAR HCE1 $\rightarrow 0.4 \text{ H2O} + 0.39 \text{ CO2} + 0.05 \text{ HCOOH} + 0.49 \text{ CO} + 0.01 \text{ G{CO}}$ | $3.0 \times 10^{-3} \times T$ | 3,600 |
| 9 | $+ 0.51 \text{ G}\{\text{CO2}\} + 0.05 \text{ G}\{\text{H2}\} + 0.4 \text{ CH2O} + 0.43 \text{ G}\{\text{COH2}\} \text{ loose} + 0.3$ | 3.0 × 10 ° × 1 | 5,000 |
| | $CH4 + 0.325 G\{CH4\} + 0.1 C2H4 + 0.075 G\{C2H4\} + 0.975 CHAR + 0.37$ | | |
| | $G\{COH2\}$ stiff + 0.1 $H2$ + 0.2 $G\{C2H4\}$ + 0.973 $GHAR$ + 0.57 | | |
| 10 | $HCE2 \rightarrow 0.3 \text{ CO} + 0.5125 \text{ CO2} + 0.1895 \text{ CH4} + 0.5505 \text{ H2} + 0.056 \text{ H2O} +$ | 7.0×10^{9} | 30,500 |
| 10 | 0.049 C2H5OH + 0.035 CH2OHCHO + 0.105 CH3CO2H + 0.0175 HCOOH | 1.0 × 10 | 30,300 |
| | $+ 0.145 \text{ FURFURAL} + 0.05 \text{ G{CH}} + 0.105 $ | | |
| | $0.45 \text{ G}\{\text{CO2}\} + 0.18 \text{ G}\{\text{COH2}\} \text{ loose} + 0.7125 \text{ CHAR} + 0.21 \text{ G}\{\text{H2}\} + 0.78 \text{ G}\{\text{CO2}\} + 0.18 \text{ G}\{\text{COH2}\} \text{ loose} + 0.7125 \text{ CHAR} + 0.21 \text{ G}\{\text{H2}\} + 0.78 \text{ G}\{\text{CO2}\} + 0.78 \text{ G}\{C$ | | |
| | G{COH2} stiff + 0.2 G{C2H6} | | |
| 11 | LIGH \rightarrow LIGOH + 0.5 C2H5CHO + 0.4 C2H4 + 0.2 CH2OHCHO + 0.1 CO | 6.7×10^{12} | 37,500 |
| | + 0.1 C2H6 | VI, II = V | 0.,000 |
| 12 | $LIGO \rightarrow LIGOH + CO2$ | 3.3×10^{8} | 25,500 |
| 13 | $LIGC \rightarrow 0.35 \ LIGCC + 0.1 \ VANILLIN + 0.1 \ C6H5OCH3 + 0.27 \ C2H4 +$ | 1.0×10^{11} | 37,200 |
| | $H2O + 0.17 G\{COH2\} loose + 0.4 G\{COH2\} stiff + 0.22 CH2O + 0.21 CO$ | | , |
| | $+ 0.1 \text{ CO2} + 0.36 \text{ G}\{\text{CH4}\} + 5.85 \text{ CHAR} + 0.2 \text{ G}\{\text{C2H6}\} + 0.1 \text{ G}\{\text{H2}\}$ | | |
| 14 | LIGCC \rightarrow 0.25 VANILLIN + 0.15 CRESOL + 0.15 C6H5OCH3 + 0.35 | 1.0×10^{4} | 24,800 |
| | CH2OHCHO + 0.7 H2O + 0.45 CH4 + 0.3 C2H4 + 0.7 H2 + 1.15 CO + | | |
| | $0.4 \text{ G}\{\text{CO}\} + 6.80 \text{ CHAR} + 0.4 \text{ C2H6}$ | | |
| 15 | LIGOH \rightarrow 0.9 LIG + H2O + 0.1 CH4 + 0.6 CH3OH + 0.3 G{CH3OH} + | 1.5×10^{8} | 30,000 |
| | 0.05 CO2 + 0.65 CO + 0.6 G(CO) + 0.05 HCOOH + 0.45 G(COH2) loose + | | |
| | $0.4 \text{ G}\{\text{COH2}\} \text{ stiff} + 0.25 \text{ G}\{\text{CH4}\} + 0.1 \text{ G}\{\text{C2H4}\} + 0.15 \text{ G}\{\text{C2H6}\} + 4.25$ | | |
| | CHAR + 0.025 C24H28O4 + 0.1 C2H3CHO | | |
| 16 | $LIG \rightarrow VANILLIN + 0.1 C6H5OCH3 + 0.5 C2H4 + 0.6 CO + 0.3 CH3CHO$ | $4.0 \times T$ | 12,000 |
| | + 0.1 CHAR | 0 | |
| 17 | $LIG \rightarrow 0.6 \text{ H2O} + 0.3 \text{ CO} + 0.1 \text{ CO2} + 0.2 \text{ CH4} + 0.4 \text{ CH2O} + 0.2 \text{ G{CO}}$ | $8.3 \times 10^{-2} \times T$ | 8,000 |
| | $+ 0.4 \text{ G}\{\text{CH4}\} + 0.5 \text{ G}\{\text{C2H4}\} + 0.4 \text{ G}\{\text{CH3OH}\} + 1.25 \text{ G}\{\text{COH2}\} \text{ loose} +$ | | |
| | $0.65 \text{ G}\{\text{COH2}\} \text{ stiff} + 6.1 \text{ CHAR} + 0.1 \text{ G}\{\text{H2}\}$ | 0 | |
| 18 | $LIG \rightarrow 0.6 \text{ H2O} + 2.6 \text{ CO} + 0.6 \text{ CH4} + 0.4 \text{ CH2O} + 0.75 \text{ C2H4} + 0.4 \text{ CH3OH}$ | 1.5×10^{9} | 31,500 |
| 4 - | + 4.5 CHAR + 0.5 C2H6 | = 0 4610 | |
| 19 | $TGL \rightarrow C2H3CHO + 2.5 MLINO + 0.5 U2ME12$ | 7.0×10^{12} | 45,700 |
| 20 | $TANN \rightarrow 0.85 C6H5OH + 0.15 G\{C6H5OH\} + G\{CO\} + H2O + ITANN$ | 2.0×10^{1} | 10,000 |
| | | | |

| 21 | $ITANN \rightarrow 5 \text{ CHAR} + 2 \text{ CO} + \text{H2O} + 0.55 \text{ G}\{\text{COH2}\} \text{ loose} + 0.45 \text{ G}\{\text{COH2}\}$ | 1.0×10^{3} | 25,000 |
|----|--|----------------------|--------|
| | stiff | | |
| 22 | $G\{CO2\} \to CO2$ | 1.0×10^{6} | 24,500 |
| 23 | $G\{CO\} \to CO$ | 5.0×10^{12} | 52,500 |
| 24 | $G\{CH3OH\} \rightarrow CH3OH$ | 2.0×10^{12} | 50,000 |
| 25 | $G{COH2}loose \rightarrow 0.2 CO + 0.2 H2 + 0.8 H2O + 0.8 CHAR$ | 6.0×10^{10} | 50,000 |
| 26 | $G\{C2H6\} \rightarrow C2H6$ | 1.0×10^{11} | 52,000 |
| 27 | $G\{CH4\} \rightarrow CH4$ | 1.0×10^{11} | 53,000 |
| 28 | $G\{C2H4\} \rightarrow C2H4$ | 1.0×10^{11} | 54,000 |
| 29 | $G\{C6H5OH\} \rightarrow C6H5OH$ | 1.5×10^{12} | 55,000 |
| 30 | $G{COH2}$ stiff $\rightarrow 0.8 CO + 0.8 H2 + 0.2 H2O + 0.2 CHAR$ | 1.0×10^{9} | 59,000 |
| 31 | $G\{H2\} 	o H2$ | 1.0×10^{8} | 70,000 |
| 32 | $ACQUA \rightarrow H2O$ | $1.0 \times T$ | 8,000 |

Table 16: Description of the chemical species in the Debiagi kinetics scheme for biomass pyrolysis. Source [4].

| Item | Name | Formula | Phase | Description |
|------|-------------------|----------------------|-------------|----------------------------|
| 1 | CELL | $C_6H_{10}O_5$ | solid | cellulose |
| 2 | CELLA | $C_6H_{10}O_5$ | solid | active cellulose |
| 3 | GMSW | $C_5H_8O_4$ | solid | hemicellulose softwood |
| 4 | XYHW | $C_5H_8O_4$ | solid | hemicellulose hardwood |
| 5 | XYGR | $C_5H_8O_4$ | solid | hemicellulose grass |
| 6 | HCE1 | $C_5H_8O_4$ | solid | intermediate hemicellulose |
| 7 | HCE2 | $C_5H_8O_4$ | solid | intermediate hemicellulose |
| 8 | ITANN | $C_8H_4O_4$ | solid | intermediate phenolics |
| 9 | LIG | $C_{11}H_{12}O_4$ | solid | intermediate lignin |
| 10 | LIGC | $C_{15}H_{14}O_4$ | solid | carbon rich lignin |
| 11 | LIGCC | $C_{15}H_{14}O_4$ | solid | intermediate lignin |
| 12 | LIGH | $C_{22}H_{28}O_9$ | solid | hydrogen rich lignin |
| 13 | LIGO | $C_{20}H_{22}O_{10}$ | solid | oxygen rich lignin |
| 14 | LIGOH | $C_{19}H_{22}O_8$ | solid | intermediate lignin |
| 15 | TANN | $C_{15}H_{12}O_7$ | solid | tannins |
| 16 | TGL | $C_{57}H_{100}O_{7}$ | solid | triglycerides |
| 17 | CHAR | $^{\mathrm{C}}$ | solid | char as pure carbon |
| 18 | $G\{COH2\}$ loose | CH_2O | metaplastic | loose formaldehyde |
| 19 | $G\{CO2\}$ | CO_2 | metaplastic | trapped carbon dioxide |
| 20 | $G\{CO\}$ | CO | metaplastic | trapped carbon monoxide |
| 21 | $G\{CH3OH\}$ | CH_4O | metaplastic | trapped methanol |
| 22 | $G\{CH4\}$ | CH_4 | metaplastic | trapped methane |
| 23 | $G\{C2H4\}$ | C_2H_4 | metaplastic | trapped ethylene |
| 24 | $G\{C6H5OH\}$ | C_6H_6O | metaplastic | trapped phenol |
| 25 | $G\{COH2\}$ stiff | CH_2O | metaplastic | stiff formaldehyde |
| 26 | $G{H2}$ | H_2 | metaplastic | trapped hydrogen |
| 27 | $G\{C2H6\}$ | C_2H_6 | metaplastic | trapped ethane |
| 28 | C2H4 | C_2H_4 | gas | ethylene |
| 29 | C2H6 | C_2H_6 | gas | ethane |
| 30 | CH2O | $\mathrm{CH_{2}O}$ | gas | formaldehyde |
| 31 | CH4 | CH_4 | gas | methane |
| 32 | CO | CO | gas | carbon monoxide |
| 33 | CO2 | CO_2 | gas | carbon dioxide |
| 34 | H2 | H_2 | gas | hydrogen |
| 35 | C2H3CHO | C_3H_4O | liquid | acrolein |

| 36 | C2H5CHO | C_3H_6O | liquid | propionaldehyde |
|----|-------------|---------------------|--------|-------------------------------|
| 37 | C2H5OH | C_2H_6O | liquid | ethanol |
| 38 | C5H8O4 | $C_5H_8O_4$ | liquid | xylofuranose |
| 39 | C6H10O5 | $C_6H_{10}O_5$ | liquid | levoglucosan |
| 40 | C6H5OCH3 | C_7H_8O | liquid | anisole |
| 41 | C6H5OH | C_6H_6O | liquid | phenol |
| 42 | C6H6O3 | $C_6H_6O_3$ | liquid | hydroxymethylfurfural |
| 43 | C24H28O4 | $C_{24}H_{28}O_4$ | liquid | heavy molecular weight lignin |
| 44 | CH2OHCH2CHO | $C_3H_6O_2$ | liquid | propionic acid |
| 45 | CH2OHCHO | $C_2H_4O_2$ | liquid | acetic acid |
| 46 | СНЗСНО | C_2H_4O | liquid | acetaldehyde |
| 47 | CH3CO2H | $C_2H_4O_2$ | liquid | acetic acid |
| 48 | CH3OH | $\mathrm{CH_{4}O}$ | liquid | methanol |
| 49 | CHOCHO | $C_2H_2O_2$ | liquid | glyoxal |
| 50 | CRESOL | C_7H_8O | liquid | cresol |
| 51 | FURFURAL | $C_5H_4O_2$ | liquid | 2-furaldehyde |
| 52 | H2O | H_2O | liquid | water from reactions |
| 53 | HCOOH | CH_2O_2 | liquid | formic acid |
| 54 | MLINO | $C_{19}H_{34}O_{2}$ | liquid | methyl linoleate |
| 55 | U2ME12 | $C_{13}H_{22}O_2$ | liquid | linalyl propionate |
| 56 | VANILLIN | $C_8H_8O_3$ | liquid | vanillin |
| 57 | ACQUA | H_2O | liquid | water within biomass |
| | | | | |

3.2 Biomass characterization

The Debiagi kinetics rely on an initial biomass composition defined as cellulolose, hemicellulose, lignin-c, lignin-h, lignin-o, tannins, and triglycerides. According to the Debiagi et al. 2015 paper, the chemical components of the biomass are defined as shown below in Table 17 [5]. The Debiagi article does not provide information on how to experimentally determine these components; therefore, the reader must decide on appropriate measurement techniques.

Table 17: Chemical components of biomass according to Debiagi et al. [5].

| Biomass composition | Description |
|---------------------|--|
| cellulose | glucan |
| hemicellulose | mixture of sugars such as hexoses and pentoses; mainly xylose, man- nose, galactose, and arabinose |
| lignin | aromatic alcohols such as coniferyl, sinapyl, p-coumaryl alcohol |
| lignin-c | carbon-rich lignin |
| lignin-h | hydrogen-rich lignin |
| lignin-o | oxygen-rich lignin |
| tannins | hydrophilic extractives, phenolics, ethanol and water, represented by a gallocatechin polymer |
| triglycerides | hydrophobic extractives, hexane and ether, linoleic acid |

Ideally, the composition of the biomass would be directly measured; otherwise, the characterization method discussed in the Debiagi paper estimates the composition based on elemental analysis data [5]. The characterization method utilizes carbon and hydrogen obtained from elemental (ultimate) analysis of the biomass to predict the biochemical composition in terms of cellulose, hemicellulose, and lignin. Splitting parameters α , β , γ , δ , ϵ are used to improve the validity of the characterization procedure by accounting for extractives in the biomass.

3.3 Batch reactor

The material balance for a typical chemical reactor is shown in Equation 1 where C_0 is inlet concentration, C is outlet concentration, v is volumetric flow rate, r is the reaction rate, and V is the reactor volume.

$$accumulation = input - output + reaction$$

$$\frac{dC}{dt}V = vC_0 - vC + rV$$
(1)

A batch reactor was modeled to understand the time scales associated with the biomass pyrolysis kinetics. For the batch reactor, input and output is zero therefore only the accumulation and reaction terms remain in the material balance. For a constant volumne reactor the V terms cancel out; therefore, Equation 2 represents the material balance for a batch reactor model.

$$accumulation = 0 - 0 + reaction$$

$$\frac{dC}{dt} = r$$
(2)

3.4 Sensitivity analysis

A sensitivity analysis was performed with the Debiagi pyrolysis kinetics to investigate the effects of biomass composition on product yields. The awesome SALib Python package was utilized for sample generation and prediction of the Sobol indices [6]. For the sensitivity analysis model, a sample represents the biomass composition as cellulose, hemicellulose, lignin-c, lignin-h, lignin-o, tannins, and triglycerides. This sample (or composition) is used in a reactor model at a certain temperature and pressure to predict pyrolysis yields. The sensitivity analysis model applies this approach to a large sample matrix then uses the generated data to perform a Sobol analysis.

4 Results and discussion

Characterization of the biomass along with batch reactor and sensitivity analysis results are discussed in this section.

4.1 Pyrolysis kinetics sensitivity analysis

A sensitivity analysis of the Debiagi pyrolysis kinetics in a batch reactor model was performed using 16,000 samples, a reaction time of 10 seconds, a reactor temperature of 773.15 K, and a reactor pressure of 101,325 Pa. Each sample represents some combination of cellulose, hemicellulose, lignin-c, lignin-h, lignin-o, tannins, and triglycerides. Batch reactor yields from the samples are shown in Figures 5 and 6. The Sobol sensitivity analysis based on the batch reactor results is presented in Figure 7.

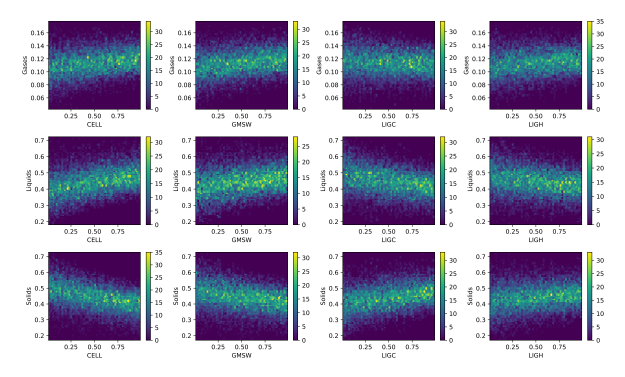


Figure 5: Batch reactor results for cellulose, hemicellulose (GMSW), carbon-rich lignin (LIGC), and hydrogen-rich lignin (LIGH) using 16,000 samples. Reaction time is 10 seconds at 773.15 K and 101,325 Pa. Colorbar represents bin count.

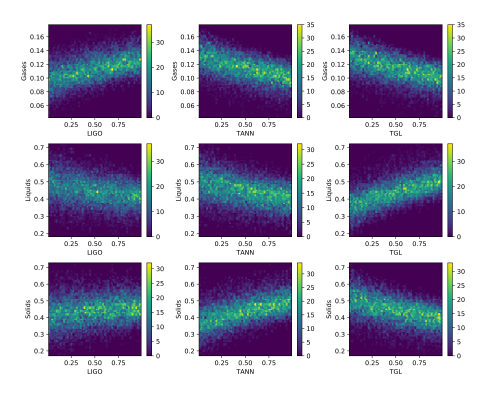


Figure 6: Batch reactor results for oxygen-rich lignin (LIGO), tannins (TANN), and triglycerides (TGL) using 16,000 samples. Reaction time is 10 seconds at 773.15 K and 101,325 Pa. Colorbar represents bin count.

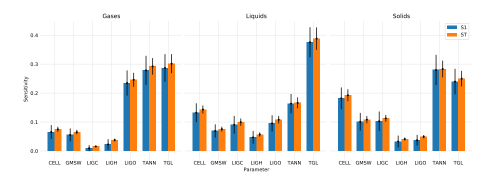


Figure 7: First-order (S1) and total-order (ST) Sobol indices for biomass composition with reactants grouped as gases, liquids, and solids using 16,000 samples.

4.2 Blend3 biomass composition

Several approaches were investigated to characterize the Blend3 feedstock for use with the Debiagi pyrolysis kinetics. The first approach uses the characterization method discussed in the Debiagi et al. 2015 paper where carbon and hydrogen from ultimate analysis is used to determine the biomass composition [5]. To use this approach for the Blend3 feedstock, the mass fraction of C and H on a dry ash-free basis (last column in Table 18) is used for the biomass characterization procedure.

Table 18: Ultimate analysis bases for the Blend3 feedstock. Mass percent values are given for as-received (ar), dry, and dry ash-free (daf) basis.

| Element | % ar | % dry | % daf | % daf |
|--------------|-------|-------|-------|-------|
| С | 49.52 | 52.70 | 53.06 | 53.16 |
| \mathbf{H} | 5.28 | 5.62 | 5.66 | 5.67 |
| O | 38.35 | 40.82 | 41.10 | 41.17 |
| N | 0.15 | 0.16 | 0.16 | |
| S | 0.02 | 0.02 | 0.02 | |
| ash | 0.64 | 0.68 | | |
| moisture | 6.04 | | | |

Case 1: The first approach to characterize the Blend3 feedstock was performed using a carbon mass fraction of 53.16%, hydrogen mass fraction of 5.67%, and splitting parameters $\alpha=0.6,\ \beta=0.8,\ \gamma=0.8,\ \delta=1.0,$ and $\epsilon=1.0$ which do not account for extractives in the feedstock. Results from this characterization are shown in Figure 8 and the associated biomass composition is given in Table 20. While this approach is useful for limited feedstock data, its accuracy is questionable when compared to experimental measurements. For example, chemical analysis of the Blend3 feedstock provides a lignin composition of 29.48% (see Table 4) whereas the characterization method using ultimate analysis data estimates a total lignin composition greater than 59%.

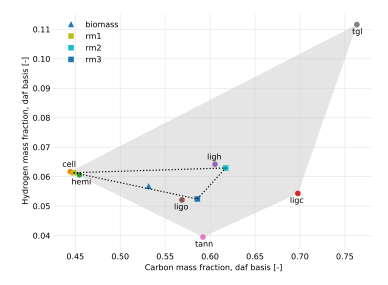


Figure 8: Characterization of the Blend3 feedstock using ultimate analysis data. Reference mixtures (rm) are labeled with square markers.

Case 2: To improve the Blend3 characterization based on ultimate analysis data, the splitting parameters were adjusted to account for extractives in the feedstock by using $\alpha=0.56,\ \beta=0.6,\ \gamma=0.6,\ \delta=0.78,$ and $\epsilon=0.88.$ Also, since the uncertainty in the ultimate analysis data is unknown (see Table 3) the carbon mass fraction was adjusted to 51% and the hydrogen mass fraction to 6%. Results from these adjustments are presented in Figure 9 and Table 20.

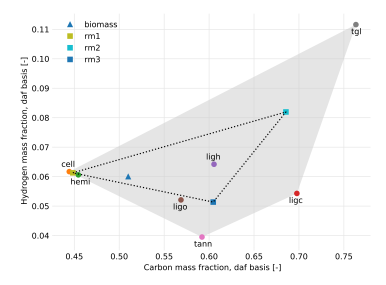


Figure 9: Characterization of the Blend3 feedstock using modified ultimate analysis data and adjusting the splitting parameters to account for extractives. Reference mixtures (rm) are labeled with square markers.

Case 3: The final approach to characterize the Blend3 feedstock, was to use chemical analysis data (see Table 4) to determine the biomass composition. As summarized in Table 19 the cellulose is represented by glucan while hemicellulose is comprised of xylan, mannan, galactan, arabinan, free fructose, free glucose, and sucrose. The measurement technique to determine the lignin components is unknown; therefore, the lignin is evenly divided into the carbon, hydrogen, and oxygen-rich fractions. Tannins are represented by acetyl, water extractives, and ethanol extractives while ash is the non-structural and structural inorganics. Triglycerides were not present in the Blend3 feedstock. Finally, the biomass composition based on the chemical analysis measurements is given in Table 20.

Table 19: Biomass composition represented by chemical analysis components.

| Biomass composition | Chemical analysis |
|---------------------|---|
| cellulose | glucan |
| hemicullulose | xylan, mannan, galactan, arabinan, free fructose, free glucose, sucrose |
| lignin-c | lignin / 3 |
| lignin-h | lignin / 3 |
| lignin-o | lignin / 3 |
| tannins | acetyl, water extractives, ethanol extractives |
| triglycerides | ? |
| ash | non-structural inorganics, structural inorganics |

Table 20: Biomass composition for the Blend3 feedstock. Values are reported as mass percent on a dry ash-free basis (% daf).

| Biomass composition | Case 1 | Case 2 | Case 3 |
|---------------------|--------|--------|--------|
| cellulose | 26.38 | 39.24 | 39.19 |
| hemicellulose | 14.33 | 25.12 | 23.26 |
| lignin-c | 7.84 | 8.57 | 9.89 |
| lignin-h | 5.27 | 3.11 | 9.89 |
| lignin-o | 46.18 | 18.00 | 9.89 |
| tannins | 0.00 | 2.95 | 7.88 |
| triglycerides | 0.00 | 3.01 | 0.00 |

4.3 Blend3 batch reactor conversion and yields

Here.

5 Conclusions

Here.

6 Source code

The Python code used to develop the models and generate results discussed in this paper is available on GitHub at https://github.com/ccpcode/nrel-efr.

7 Computational resources

An Apple MacBook Pro laptop was used to develop all the models and generate all the results discussed in this paper. A summary of the hardware is listed below:

• Model: MacBook Pro (16-inch, 2019)

 \bullet Processor: 2.6 GHz 6-Core Intel i7

• Memory: 32 GB 2667 MHz DDR4

• Integrated Graphics: Intel UHD Graphics 630

• Discrete Graphics: 4GB AMD Radeon Pro 5500M

A Appendix

A.1 Sensitivity analysis

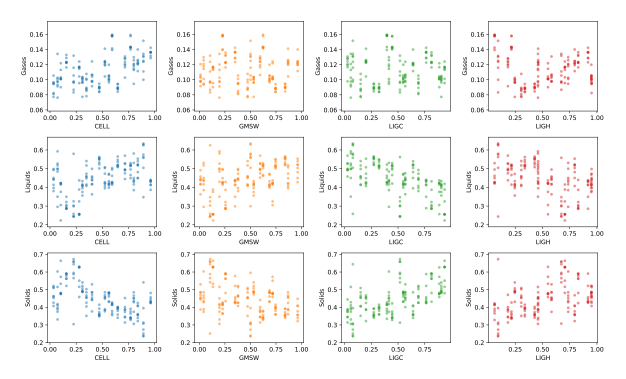


Figure 10: Batch reactor results for cellulose, hemicellulose (GMSW), carbon-rich lignin (LIGC), and hydrogen-rich lignin (LIGH) using 160 samples. Reaction time is 10 seconds at 773.15 K and 101,325 Pa.

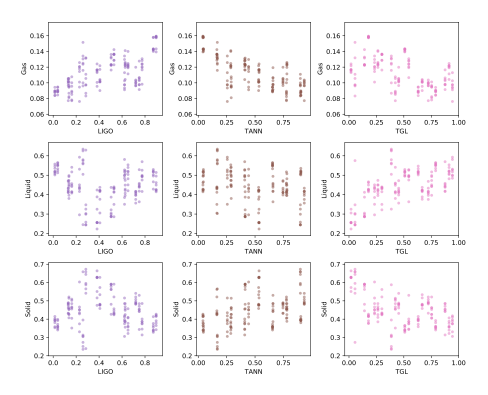


Figure 11: Batch reactor results for oxygen-rich lignin (LIGO), tannins (TANN), and triglycerides (TGL) using 160 samples. Reaction time is 10 seconds at $773.15~\mathrm{K}$ and $101,325~\mathrm{Pa}$.

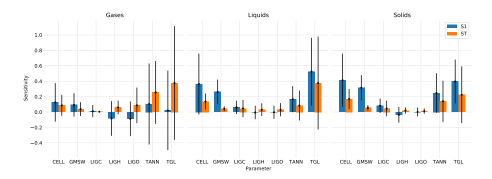


Figure 12: First-order (S1) and total-order (ST) Sobol indices for biomass composition with reactants grouped as gases, liquids, and solids using 160 samples.

References

- [1] Unknown Author. *CHN Analysis Report*. Tech. rep. From Excel spreadsheet jw-pine-chn-report-jw190920. Unknown institution, 2019.
- [2] Unknown Author. Summary normalized. Tech. rep. From Excel spreadsheet summary-normalized. Unknown institution, 2020.
- [3] Mike Choratch. *Proximate and ultimate analysis*. Tech. rep. From PDF chns_ipc_ms_blend3_nrel2017. Hazen Research, Inc, 2017.
- [4] P. Debiagi et al. "A predictive model of biochar formation and characterization". In: *Journal of Analytical and Applied Pyrolysis* 134 (2018), pp. 326–335.
- [5] Paulo Eduardo Amaral Debiagi et al. "Extractives Extend the Applicability of Multistep Kinetic Scheme of Biomass Pyrolysis". In: *Energy & Fuels* 29.10 (2015), pp. 6544–6555.
- [6] Jon Herman and Will Usher. "SALib: An open-source Python library for Sensitivity Analysis". In: Journal of Open Source Software 2.9 (2017), p. 97. DOI: 10.21105/joss.00097. URL: https://doi.org/10.21105/joss. 00097.
- [7] M. Brennan Pecha et al. "Integrated Particle- and Reactor-Scale Simulation of Pine Pyrolysis in a Fluidized Bed". In: *Energy & Fuels* 32.10 (2018), pp. 10683–10694.
- [8] Anne Starace and Justin Sluiter. Compositional analysis Blend3. Tech. rep. From Excel spreadsheet comp_analysis_blend3_fy17verification. National Renewable Energy Laboratory, 2020.