Toward net-zero sustainable aviation fuel with wet waste–derived volatile fatty acids

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With the increasing demand for net-zero sustainable aviation fuels (SAF), new conversion technologies are needed to process waste feedstocks and meet carbon reduction and cost targets. Wet waste is a low-cost, prevalent feedstock with the energy potential to displace over 20% of US jet fuel consumption; however, its complexity and high moisture typically regulates its use to methane production from anaerobic digestion. To overcome this, methanogenesis can be arrested during fermentation to instead produce C2 to C4 volatile fatty acids (VFA) for catalytic upgrading to SAF. Here, we evaluate the catalytic conversion of food waste–derived VFAs to produce n-paraffin SAF for near-term use as a 10 vol% blend for ASTM “Fast Track” qualification and produce a highly branched, isoparaffin VFA-SAF to increase the renewable blend limit. VFA ketonization models assessed the carbon chain length distributions suitable for each VFA-SAF conversion pathway, and food waste–derived VFA ketonization was demonstrated for >100 h of time on stream at approximately theoretical yield. Fuel property blending models and experimental testing determined normal paraffin VFA-SAF meets 10 vol% fuel specifications for “Fast Track.” Synergistic blending with isoparaffin VFA-SAF increased the blend limit to 70 vol% by addressing flashpoint and viscosity constraints, with sooting 34% lower than fossil jet. Techno-economic analysis evaluated the major catalytic process cost-drivers, determining the minimum fuel selling price as a function of VFA production for SAF that avoid direct competition with food resources. Life cycle analysis determined that if food waste is diverted from landfills to avoid methane emissions, VFA-SAF could enable up to 165% reduction in greenhouse gas emissions relative to fossil jet.

Significance

To meet the growing demand for sustainable aviation fuels (SAF), conversion pathways are needed that leverage waste carbon and meet jet fuel property specifications. Here, we demonstrate SAF production from food waste–derived volatile fatty acids (VFA) by targeting normal paraffins for a near-term path to market and branched isoparaffins to increase the renewable content long term. Combining these distinct paraffin structures was shown to synergistically improve VFA-SAF flash point and viscosity to increase the renewable blend limit to 70%. Life cycle analysis shows the dramatic impact on the carbon footprint if food waste is diverted from landfills to produce VFA-SAF, highlighting the potential to meet jet fuel safety, operability, and environmental goals.

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Competing interest statement: N.A.H., X.H., H.N., and D.R.V. are inventors on a patent application submitted by the Alliance for Sustainable Energy on methods for the production of VFA-SAF (US 17/121,336 filed on December 14, 2020).

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waste (2.5 BPGY), animal manure (4.4 BPGY), wastewater sludge (1.9 BPGY), and the abovementioned waste fats, oils, and grease (1.7 BPGY) (14, 15). While waste lipid feedstocks may be best suited for HEFA refining, valorization strategies are needed for the remaining wet waste feedstocks. Diverting food waste from landfills is of particular note for reducing greenhouse gas emissions, as landfilling one dry ton of food waste has been estimated to release as much as 1.8 tons of CO₂ equivalents, assuming landfill methane is collected and recovered for electricity generation (16, 17). Globally, food waste accounts for 6% of greenhouse emissions (18). The high moisture content of wet waste restricts the use of conventional thermochemical conversion approaches (e.g., pyrolysis and gasification) used to produce liquid biofuels from terrestrial biomass, directing technology development efforts toward hydrothermal liquefaction, biological conversion, and hybrid processes (19).

Currently, anaerobic digestion to produce biogas is the leading technology to recover energy from wet waste (20). The high moisture content of wet waste limits its transport and necessitates local processing, with the majority of US anaerobic digestion facilities located near population-dense areas and airports (21). Biogas purification provides a route to pipeline quality renewable natural gas compatible with existing infrastructure. Life cycle analysis has shown that negative carbon intensity can be achieved when producing renewable natural gas from municipal solid waste (~23 g CO₂eq/MJ) and dairy waste (~276 g CO₂eq/MJ), providing a significant economic driver under the LCFS (12). While renewable natural gas targets an enormous US market (~246 BGPY of jet fuel equivalent) (10), producing liquid hydrocarbon fuels from wet waste offers the potential to address the challenge of decarbonizing the aviation sector.

Anaerobic digestion of wet waste can be arrested prior to methanogenesis to generate both short chain (C₂ to C₆) and medium chain length (C₆ to C₈) carboxylic acids as precursors for biofuels and biobased chemicals (14, 22–27), hereon collectively referred to as volatile fatty acids (VFAs). VFA production by arrested methanogenesis offers the potential to utilize existing biogas infrastructure and a wide variety of wet waste feedstocks (14, 22, 28) with ongoing research and development working to increase VFA titers, rates, and yields by tailoring feedstock composition, microbial consortia, fermentation parameters, and online separation technologies (14, 22, 29–31). Currently, C₂ to C₆ carboxylic acids are primarily produced from the oxidation of petroleum derivatives, while C₆ and C₈ carboxylic acids are primarily derived from coconut and palm oil (29). Propionic acid (C₃) and butyric acid (C₄) address chemical market volumes on the order of 0.1 to 0.2 BGPY (29), while medium chain length carboxylic acids target specialty markets. Given the availability of wet waste and potential saturation of biobased chemical markets in the long term, VFAs provide a potential target intermediate for catalytic upgrading into low carbon intensity biofuel (23, 25, 26, 32–35).

VFAs can be catalytically upgraded to SAF through carbon coupling and deoxygenation chemistries. Depending on their chain length, VFAs can be converted into normal paraffins identical to those found in petroleum or undergo an additional carbon coupling step to generate isoparaffin, cycloparaffin, and aromatic hydrocarbons with molecular structures distinct from fossil jet (Fig. 1).

Ketonization is the first unit operation to elongate the carbon backbone of VFAs (14, 22). Ketonization reacts two VFAs to produce a single ketone that is one carbon shorter than the sum of both acids and removes oxygen in the form of water and carbon dioxide (36, 37). Ketonization of acetic acid to acetone has been commercialized (37), with longer chain acids actively researched for biofuel and biochemical applications. Following ketonization, ketones ≥C₈ can undergo direct hydrodeoxygenation (Fig. 1, Top) to produce normal paraffin-rich hydrocarbons. In comparison, ketones ≤C₇ require a second carbon coupling step prior to hydrodeoxygenation to fall within the C₆ to C₁₈ range of jet fuel (Fig. 1, Bottom). Ketone carbon coupling can take place by various pathways including aldol condensation chemistry (25) as well as ketone reduction to alcohols for further dehydration and oligomerization (32, 34). Aldol condensation of central ketones is an emerging bench-scale chemistry that can generate structurally unique isoparaffins (38) with significantly lower freezing points for jet fuel applications due to the high degree of branching as well as reduce the intrinsic sooting tendency relative to aromatic hydrocarbons by over twofold (25).

Normal paraffins produced from VFAs (Fig. 1, Top) can provide fungible hydrocarbons identical to those in petroleum that offer a near-term path to SAF qualification and market entry. In the United States, new SAF conversion routes must complete a rigorous qualification process to ensure fuel safety and operability overseen by ASTM International, the Federal Aviation Administration, and aviation original equipment manufacturers (OEMs). There are currently seven ASTM-approved routes for SAF that are derived from Fischer–Tropsch processing of syngas, the abovementioned esters and fatty acids, farne- sene, ethanol, isobutanol, and algal hydrocarbons (39). Further details summarizing current ASTM-qualified routes to SAF can be found in SI Appendix, Table S1. Historically, ASTM qualification can require jet fuel volumes on order of over 100,000 gallons in order to pass a four-tiered screening process and two OEM review stage gates that may take place over a period of 3 to 6 years.

Fig. 1. Overview scheme of the major oxygenate and hydrocarbon molecules produced when converting wet waste VFA into Fast Track VFA-SAF that is composed of normal paraffin-rich hydrocarbons (Top Right) and Aldol Condensation VFA-SAF composed of isoparaffin-rich hydrocarbons (Bottom Right).
Results and Discussion

A simplified kinetic model was developed and used to evaluate carbon flow to the suitable ketone upgrading pathways based on the incoming VFA chain length distribution, as shown in Fig. 2A and D. Both the C6/C8 and C4/C6 VFA sample carbon chain length distributions were modeled for accounting for higher rates of cross-ketonization relative to self-ketonization (47). Model results were then compared against ketone profiles determined experimentally under complete conversion conditions (Fig. 2C and F and SI Appendix, Figs. S1–S3). Modeled ketone chain length distributions were in good agreement with experimental results, with <10% difference in the mole fraction for a given carbon number. The C4/C6 VFA sample yielded a majority of ketones with chain lengths ≥C8 suitable for direct hydrodeoxygenation for the Fast Track VFA-SAF conversion pathway (95% modeled and 95% experimental). In comparison, slightly less than half of the ketones produced from the C6/C8 VFA sample were ≥C8 (47% modeled and 44% experimental), with a near equal portion ≤C6 suitable for the Aldol Condensation VFA-SAF conversion pathway. To demonstrate the robustness of this model, an additional VFA profile was experimentally tested and shown to be in close agreement with the predictive ketone carbon number distribution model (SI Appendix, Fig. S2).

Ketonization Catalyst Performance. The stability of a commercial ZrO2 catalyst for ketonization was evaluated with both model and biogenic VFAs derived from food waste. Previous studies by our team have shown full conversion of butyric acid with near theoretical ketone yield using this catalyst (25); therefore, partial conversion conditions were evaluated with mixed model VFAs to assess ketone stability during 72 h of TOS (SI Appendix, Fig. S4). The first 48 h of TOS resulted in 6% drop in conversion which appeared to stabilize with the final 24 h showing only 2% drop in conversion. This suggests carbon laydown may initially occur on strong acid sites (48) which may be manageable with periodic oxidative regeneration. Complete and partial conversion runs were then performed with biogenic VFAs to assess the deposition of biogenic impurities and regenerability of the catalyst. The biogenic VFA feed contained trace alkali impurities, as well as sulfur (Table 1), which may deposit on the catalyst surface and poison acid sites over time, particularly alkali elements that are known to deactivate reducible metal oxides (49, 50). Both the C6/C8 and C4/C6 VFA samples were prepared for catalytic testing for ASTMS Fast Track, hereon referred to as “Fast Track VFA-SAF.” In parallel, VFA-derived ketones ≤C7 were processed via aldol condensation and hydrodeoxygenation to produce predominantly isoparaffin hydrocarbons for Tier α and Tier β prescreening, hereon referred to as “Aldol Condensation VFA-SAF.” Higher blends with both VFA-SAF fractions were examined for increased ketone content and reduce soot formation while still meeting fuel property specifications. Lastly, techno-economic and life cycle analysis was performed to evaluate the sensitivity of catalytic process parameters on VFA-SAF production costs as well as potential greenhouse gas reductions relative to fossil jet.

VFA Ketonization Model. A simplified kinetic model was developed and used to evaluate carbon flow to the suitable ketone upgrading pathways based on the incoming VFA chain length distribution, as shown in Fig. 2A and D. Both the C6/C8 and C4/C6 VFA sample carbon chain length distributions were modeled by accounting for higher rates of cross-ketonization relative to self-ketonization (47). Model results were then compared against ketone profiles determined experimentally under complete conversion conditions (Fig. 2C and F and SI Appendix, Figs. S1–S3).

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volume decreased by <10%, consistent with the relatively low coke content. Regeneration of the 100-h spent ZrO2 catalyst exposed to biogenic impurities was performed by a typical oxidation cycle in flowing air at 500 °C. Following regeneration, carbon was no longer detected on the catalyst surface by TGA. and similar textural properties and surface acidity were observed as the fresh catalyst (Table 2 and SI Appendix, Figs. S6 and S7). Ketonization partial conversion tests (~37% VFA conversion, 290 °C, weight hourly space velocity of 7.7 h−1) confirmed comparable ketonization activity between the fresh and regenerated catalyst using the biogenic C4/C6 VFA sample with online detection of effluent gas CO2 by nondispersive infrared detection (NDIR), with differences within experimental error (Fig. 3C). Analysis of the liquid products are also provided in SI Appendix, Fig. S8. As anticipated based on the spent catalyst characterization results, impurities

![Diagram](image-url)

**Fig. 2.** Modeled and experimental VFA ketonization carbon yields. For the C6/C8 VFA feed, (A) VFA carbon chain length distribution, (B) ketone and CO2 carbon yield, and (C) model and experimental ketone carbon number profile. Similar values are reported for the C4/C6 VFA feed (D–F). Blue represents ≥C8 ketone carbon chain lengths suitable for Fast Track VFA-SAF, while orange represents ≤C7 ketones that require coupling for Aldol Condensation VFA-SAF. Values are rounded to whole number.

| ppm | Neat VFA C4/C6 | Neat ketone C4/C6 | Fast Track VFA-SAF C4/C6 | Neat VFA C6/C8 | Neat ketone C6/C8 | Fast Track VFA-SAF C6/C8 |
|-----|----------------|-------------------|--------------------------|----------------|-------------------|--------------------------|
| Al  | <0.4           | <0.4              | 1.2                      | <1             | <0.4              |                          |
| B   | <0.2           | <0.2              | 1.8                      | <1             | <0.2              |                          |
| Ca  | <0.1           | <0.1              | 0.8                      | <0.1           | <0.1              |                          |
| Fe  | 0.4            | <1                | 3.0                      | <0.1           | <0.1              |                          |
| K   | <1             | <1                | 236                      | <1             | <1                |                          |
| Mg  | <0.1           | <0.1              | 4.1                      | <0.1           | <0.1              |                          |
| Mn  | <0.1           | <0.1              | <0.2                     | <0.2           | <0.2              |                          |
| N   | 37.0           | ND                | 67.7                     | 25.9           | ND                |                          |
| Na  | 11.8           | <1                | 104.8                    | <1             | <1                |                          |
| P   | <1             | <1                | 10                      | <1             | <1                |                          |
| S   | 8.2            | 1.0               | 32                       | <1             | <1                |                          |
| Si  | <1             | <1                | <1                       | <1             | <1                |                          |
| Zn  | <0.1           | <0.1              | <0.1                     | <1             | <1                | <0.1                     |

**Table 1.** Elemental analysis of the feed and upgrading products for C4/C6 and C6/C8 food waste–derived VFA samples. ND: not determined due to volume limitations.
were not detected on the regenerated ZrO$_2$ (Fig. 3B and SI Appendix, Table S4 and Fig. S5) with comparable crystallite sizes before and after oxidative treatment (SI Appendix, Fig. S9). Future work is planned to decouple the impacts of carbon laydown and biogenic impurities as a function of catalyst surface acidity, VFA chain length, and process conditions over prolonged TOS.

**Fast Track VFA-SAF.** Ketones derived from the C$_6$/C$_8$ VFA sample were processed directly by hydrodeoxygenation to produce predominantly normal paraffins suitable for Fast Track VFA-SAF. Hydrodeoxygenation was performed with a 3% Pt/Al$_2$O$_3$ catalyst prepared in house and studied in previous work (25). Hydrocarbon production was observed for up to 48 h of TOS under complete conversion conditions with a liquid mass balance of 99% (SI Appendix, Fig. S10). It should be noted that partial conversion studies are needed to evaluate hydrodeoxygenation catalyst stability with biogenic ketones in regard to both carbon laydown and impurity deactivation. This effort is ongoing by our team and beyond the scope of this initial study. Following hydrodeoxygenation, the hydrocarbon phase was decanted, and trace amounts of C$_7$ hydrocarbons ($\leq$9.8% of sample mass) were removed by distillation prior to neat and blended fuel property analysis. The overall VFA-to-hydrocarbon carbon yield was 79% (mass yield 58%) which approached theoretical (82% carbon yield and 61% mass yield).

Neat fuel properties of Fast Track VFA-SAF produced from the C$_6$/C$_8$ VFA sample were then evaluated by Tier $\alpha$ and $\beta$ screening (44) (SI Appendix, Table S2). Values were compared to specifications for an aviation fuel containing SAF as defined by ASTM D7566 Table 1 and are shown visually in Fig. 4 against gray regions representing representative fossil jet fuel properties (51). The neat Fast Track VFA-SAF displayed an average carbon number of 11.3 (Fig. 4A) which was comparable to the average value of commercial Jet A of 11.4 (SI Appendix, Table S6). The VFA-SAF sample was predominantly normal paraffins, with 7 wt % isoparaffins and 2 wt% cycloparaffins detected. Fuel property measurements of the neat sample (SI Appendix, Tables S7 and S8) confirmed its moderately higher net heat of combustion (nHOC) than the reference fossil jet fuel (43.4 MJ/kg, Jet A 43.0 MJ/kg), with viscosity and surface tension being within the typical range of fossil jet. Normalized sooting concentration measurement for a 20 vol% blend (0.87) suggests that neat Fast Track VFA-SAF extrapolates to a 65% reduction in sooting relative to fossil Jet A (SI Appendix, Table S8). However, the high concentration of small ($\leq$C$_9$) normal paraffins resulted in flash point being below specifications (31 °C, spec 38 °C), gravimetric density being below specifications (743 kg/m$^3$, spec 775 to 840 kg/m$^3$), and boiling point distribution being below the typical range for Jet A (SI Appendix, Table S8). The limited degree of molecular branching also resulted in freezing point above spec (−27 °C, spec −40 °C).

To address neat fuel property limitations, Fast Track VFA-SAF was blended at 10 vol% with Jet A. Blending at 10 vol% resulted in all measured fuel properties being within spec (Table 3 and Fig. 4A–C). The freezing point of neat Jet A increased by 5 °C to −47 °C upon blending, staying well below the limit of −40 °C. No change was observed in the flash point of Jet A after blending due to the low concentration of volatile components. The boiling point distribution of the blend was within the typical range of fossil jet fuel (Fig. 4C). Improvements in neat Jet A fuel properties upon blending included a modest increase in specific energy density to 43.2 MJ/kg, an increase in the indicated cetane number from 48 to 52, and based on the 20 vol% measurement (SI Appendix, Table S8), an anticipated decrease of 6.5% in normalized soot concentration. Additionally, the acidity of the blend is anticipated to be 0.02 mg KOH/g (based on the measured 0.15 mg KOH/g acidity of the neat fuel) which is well within the blend maximum of 0.10 mg KOH/g. While Fast Track
limits blending to 10 vol%, additional testing determined that the 20 vol% blend still met flashpoint criteria (SI Appendix, Tables S8 and S9).

**Aldol Condensation VFA-SAF.** Highly branched isoparaffins produced from Aldol Condensation VFA-SAF were then evaluated for their neat and blended Tier α and Tier β fuel properties. Batch aldol condensation reactions were performed with the $<\text{C}_8$ ketones derived from the $\text{C}_4/\text{C}_6$ VFA sample. Ketones $<\text{C}_8$ were initially separated by fractional distillation and condensed in a batch reactor. The ketones were prepared at 20 wt% in decane as an appropriate boiling point solvent that also facilitated tracking ketone conversion by GC. Nb$_2$O$_5$ powder catalyst was selected due to its high acidity and activity for internal ketone condensation. Recent work has shown internal ketones are an order of magnitude less reactive than terminal ketones and require highly acidic metal oxides (25, 52). Single-pass ketone conversion to enones was dependent on molecular structure, varying between 12 and 100% (SI Appendix, Table S10). Ketone condensation can also form cyclic and aromatic trimers, with the mechanism reviewed elsewhere (53). Carbon loss to gas-phase products was insignificant, with liquid mass balance closure $>90\%$ after catalyst filtration. Previous efforts have demonstrated the regenerability of Nb$_2$O$_5$ for 4-heptanone condensation and recycle of solvent and unreacted ketone (25), with further work needed to evaluate continuous reactor configurations and compatible solvent formulations. Enone products were recovered by distillation and processed neat over the same 3 wt% Pt/Al$_2$O$_3$ hydrodeoxygenation catalyst (SI Appendix, Fig. S10). The recovered hydrocarbon phase was used for testing without further workup.

The neat Aldol Condensation VFA-SAF showed a higher average carbon number of 13.8 due to the coupling step that converts $\text{C}_3$ to $\text{C}_7$ ketones into predominantly $\text{C}_{10}$ to $\text{C}_{14}$ enones (Fig. 4D). The sample contained 76% isoparaffins, with 13% normal paraffins, 9% monocycloparaffins, and 1% aromatics, with ring structures likely due to trimer formation during ketone condensation (38, 53, 54). The relatively low percentage of $<\text{C}10$ hydrocarbons resulted in a higher flash point of 62 °C compared to Jet A, while the high degree of branching resulted in a freezing point of $53^\circ\text{C}$ at $43.4\text{MJ/kg}$ that was above the typical range of fossil jet. Normalized sooting concentration of the 30 vol% blend was reduced by 15% relative to fossil Jet A.

**Coblending VFA-SAF.** Synergistic blending was then evaluated by combining Fast Track and Aldol Condensation VFA-SAF due to their complementary limiting fuel properties. Samples were produced from the same $\text{C}_4/\text{C}_6$ VFA starting material using ketones $<\text{C}_8$ for Fast Track and ketones $<\text{C}_6$ for Aldol Condensation. Since flashpoint is typically dictated by the lowest boiling components in the mixture, a 20 vol% blend limit was maintained for Fast Track VFA-SAF (SI Appendix, Tables S8 and S9). In contrast, viscosity is a function of the overall hydrocarbon mixture composition; therefore, it was reasoned that the low-viscosity Fast Track normal paraffins would offset the high-viscosity Aldol Condensation isoparaffins.

As shown in Fig. 4 G–I, coblending normal 20 vol% Fast Track VFA-SAF increased the blend limit of Aldol Condensation VFA-SAF from 30 vol% to 50 vol%, resulting in an overall VFA-SAF content of 70 vol%. Fuel property measurements (Table 3) determined the 70 vol% blend displayed a reduced freezing point of $61^\circ\text{C}$ and significantly increased energy density of 43.7 MJ/kg, well above the typical range for fossil jet. The high energy density of neat $\text{C}_4/\text{C}_6$ Fast Track and $\text{C}_4/\text{C}_6$ Aldol Condensation VFA-SAF, at 44.49 and 44.1 MJ/kg, respectively, is comparable to HEFA SAF, Fischer Tropsch SPK, and alcohol-to-jet (SI Appendix, Table S7) (56, 57). The high energy density can provide petroleum refiners with the ability to blend VFA-SAF into lower energy density distillates, potentially motivating adoption. Normalized soot concentration also decreased by 34% upon blending, consistent with the blend’s reduced aromatic content of 5.8%. Although the accepted low limit for aromatic compounds is 8%, a recent study indicates that cycloalkanes can replace aromatics in terms of volume swell. The combination of 5.8% aromatic and 16% cycloparaffins may be sufficient to remain above the minimum currently required for polymer seal swell (58), although further work is needed to test VFA-SAF. Cetane number of the blend increased well above the range of conventional jet fuel (CN of 64), while surface tension decreased below the typical range. The latter may have positive implications on ignition quality due to improved (decreased droplet size) fuel spray properties (59).

**Techno-Economic Analysis.** Preliminary techno-economic analysis identified the catalytic process cost drivers to guide future research for VFA-SAF upgrading. The Fast Track VFA-SAF pathway was chosen due to the higher technology and fuel readiness level relative to the Aldol Condensation pathway, since the former employs catalytic packed bed reactors for both ketonization and hydrodeoxygenation. Previous techno-economic analysis work on the aldol condensation pathway can be found elsewhere which focuses on butyric acid from corn stover hydrolysate sugar fermentation (60). The baseline process model scenario (Fig. 5A) considered the upgrading process in isolation and utilized a VFA profile corresponding to the $\text{C}_6/\text{C}_8$ sample as the starting feedstock (carbon distribution in SI Appendix, Table S3), as shorter chain VFAs are currently accessible from a wider array of waste.

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### Table 2. Fresh, spent, and regenerated ZrO$_2$ catalyst material properties used for 100 h of continuous time on stream ketonization of the $\text{C}_6/\text{C}_8$ VFA sample (Fig. 2)

| ZrO$_2$ catalyst sample | Surface area (m$^2$/g) | Pore volume (mL/g) | Total acidity (μmol/g) | Carbon content (wt%) |
|-------------------------|-----------------------|------------------|------------------------|----------------------|
| Fresh                   | 51.3                  | 0.29             | 246                    | NA                   |
| 100 h spent             | 47.6                  | 0.27             | ND                     | 1.8                  |
| Regenerated             | 48.4                  | 0.30             | 233                    | 0.0                  |

NA: not applicable for fresh catalyst. ND: not determined due to carbon laydown that interferes with measurement.
Fig. 4. VFA-SAF fuel properties relative to fossil Jet A are shown, prescreened for composition, chemical and physical properties, and distillation analysis. The first row shows Fast Track VFA-SAF blend produced from the C6/C8 VFA sample (A) neat carbon distribution, (B) 10 vol% blend fuel properties ($\sigma$ = surface tension, $\rho$ = density, $\mu$ = viscosity, $n$HOC = net heat of combustion, and CN = cetane number), and (C) 10 vol% blend simulated distillation curves. Equivalent data are shown for the Aldol Condensation VFA-SAF produced from the C4/C6 VFA sample (D–F). The 20%/50% Fast Track/Aldol Condensation blend result is shown for all three of its panels (G–I). The gray region represents the range of conventional fuels, namely, POSFs 10325, 10264, and 10289. The red regions represent out-of-specification ranges. The carbon distribution for representative fossil jet POSF 10325 is tabulated in SI Appendix, Table S6, with fuel properties in SI Appendix, Table S8. Predicted data are shown with an open circle.

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feedstocks and microbial consortia (22, 61, 62). Consistent with the VFA samples evaluated in this work, VFAs were assumed to be recovered neat from fermentation media in their nonsalt form with no additional pretreatment steps required. The VFA upgrading plant scale was based on an assumed food waste availability of 225 wet tons per day and a yield of 0.45 kg VFA/kg dry food waste. VFA ketonization yields were based on experimental results with the full suite of ketones sent for hydrodeoxygenation. Ketones ≤C7 were converted to naphtha suitable for gasoline applications. This resulted in a VFA catalytic upgrading plant capacity of 2.17 MGPY of VFA yielding 1.63 MGPY of liquid hydrocarbon biofuel (gasoline gallon equivalent energy basis). Further information on the techno-economic model and assumptions can be found in SI Appendix.

The carbon yield of VFA to fuel for the upgrading process was 85%, with 46% carbon yield to SAF and 39% to naphtha; carbon losses are attributed to the CO2 produced in the ketonization reactions and ketones lost with the aqueous phase during separation. This compares favorably to the estimated HEFA-SAF carbon yield with lipids to fuel of ∼89% (34% SAF, 50% renewable diesel, and 5% naphtha) (11). Conversely, the upstream yield of 0.45 kg VFA/kg dry food waste corresponded to an estimated carbon yield of 56%. The higher yield losses observed in the upstream food waste conversion process are mainly attributed to the CO2 and digestate resulting from anaerobic digestion. Considering the full process of food waste to fuel, an overall carbon yield of 48% was observed, with VFA-SAF and naphtha carbon yields of 26% and 22%, respectively. Further work is needed to evaluate overall process yields with additional wet waste feedstocks (e.g., manure and wastewater sludge).

With VFA production still in the early stages of development and commercialization, the cost of VFA biofuel was evaluated as a function of VFA selling price (Fig. 5B). VFA production costs are highly dependent on variables such as the food waste composition, anaerobic digestion process parameters, separation technology, and potential to sell a portion of VFAs into chemical markets. Research is ongoing to advance upstream VFA fermentation and separation technology to reduce process costs and enable the use of diverse wet waste feedstocks (14, 22, 29–31); therefore, techno-economic analysis of upstream VFA production costs was deemed to be outside the scope of this work. At a jet fuel selling price of $2.50/gallon without carbon intensity reduction credits, profitable VFA-SAF production can be realized at a maximum VFA feedstock production cost $1.08/gallon ($0.30/kg). This price point for VFA is comparable to recent lignocellulosic sugar production cost estimates from mixed feedstocks that range from $0.36 to 0.50/kg (61). Recent work has estimated that VFAs could be produced from brown algae via anaerobic digestion and membrane distillation at a cost of $0.38/kg (63). Current market prices of VFAs vary by chain length and can range from $0.40 to 2.50/kg (22); however, it should be emphasized that these market prices are based on current (largely fossil-based) production methods for individual VFAs and do not necessarily relate to production cost of mixed VFAs from a feedstock such as food waste.

Single-point sensitivity analysis was then performed on the selected downstream catalytic conversion parameters (Fig. 5C). The VFA upgrading process is largely a capital-driven process, with a capital intensity of $5.40 per gasoline gallon equivalent (GGE) of annual plant capacity. Capital estimates done here are selected downstream catalytic conversion parameters (Fig. 5C). The capital intensity of $5.40 per gasoline gallon equivalent (GGE) of annual plant capacity. Capital estimates done here are only a feasibility-level estimate and may vary significantly from actual capital expenditures. To account for this, a contingency of ±50% is applied to the overall installed plant costs. Separately, the same factor is applied to the ketonization and hydrodeoxygenation reactors individually, given that they are the largest equipment cost drivers. A 50% increase in the overall capital costs increases the minimum fuel selling price by 14%, and similar changes to the ketonization and HDO reactor costs results in minimum fuel selling price increases of 5% and 3%, respectively. The sensitivity to income tax rate was also considered, demonstrating a rate of 35% (corresponding to the US corporate tax rate prior to 2018) only resulted in a 2% increase in minimum fuel selling price compared to the base case tax rate of 21%.

The relatively low catalyst requirements of both reactors resulted in the maximum VFA production cost being rather invariant to large changes (±50%) in weight hourly space velocity and single-purchase catalyst material costs. Selectivity through each reaction was shown to have a strong impact when nontarget products were sent to the steam boiler for heat recovery, rather than contributing to biofuel. The high single-purchase cost of the...
Life Cycle Analysis. The carbon intensity of the VFA-SAF Fast Track process was then evaluated using life cycle analysis. Upstream VFA production and recovery flow rates for life cycle inventory were based on technical consultation with Earth Energy Renewables for this early-stage technology. VFA fermentation and recovery is an emerging technology area, with process flow streams anticipated to vary based on the upstream process technology configuration, waste feedstock composition, technology readiness level, and scale of implementation. VFA catalytic upgrading parameters were based on this study’s process model.

The carbon intensity for VFA-SAF was largely driven by avoided methane emissions associated with food waste, with a breakdown of associated CO2 emissions and credits shown in SI Appendix, Fig. S12. Diverting food waste from landfills resulted in negative carbon emissions by avoiding methane release (~154 g CO2eq/MJ), while VFA production and VFA catalytic upgrading accounted for 84 g CO2eq/MJ and 15 g CO2eq/MJ in emissions, respectively. This resulted in an overall carbon footprint for Fast Track VFA-SAF of ~55 g CO2eq/MJ, which is 165% lower than fossil jet fuel (85 g CO2eq/MJ). Biogenic CO2 emitted from VFA-SAF combustion was accounted for as a credit per life cycle analysis convention (69).

Using California’s LCFS credit calculator for renewable diesel, the carbon intensity reduction would provide a credit of $3.71/gallon (70). Assuming the target VFA-SAF minimum full selling price remains fixed at $2.50/gallon, this credit would allow VFA production costs to be no more than $0.71/kg (Fig. 5B). Renewable Identification Number credits under the US Environmental Protection Agency (EPA) Renewable Fuel Standard would offer additional economic incentives (71). If the 70% blend limit for VFA-SAF can be achieved with comparable life cycle inputs, the 165% reduction in lifecycle greenhouse gas emissions would provide a path toward net zero jet fuel.

While this preliminary analysis suggests the potential for significant greenhouse gas savings with VFA-SAF, it should be noted that baseline practices for wet waste management will continue to evolve, as will VFA-SAF technology. New legislation in California recently mandated a 75% reduction of landfill organics from 2014 levels by 2025 using alternative disposal practices such as composting and anaerobic digestion (72). Eliminating the avoided methane credit would lead to a carbon intensity increase by 16% compared to fossil jet fuel based on the parameters used for this analysis. Widespread changes to landfill organic waste management would impact the baseline for avoided landfill methane emission credits evaluated here and motivate VFA production from additional wet waste feedstocks including manure, wastewater sludge, algae, and lignocellulosic biomass (14, 22). In addition, advancements in renewable electricity and green hydrogen production stand to reduce the overall carbon footprint of upstream...
and downstream VFA-SAF unit operations. Significant potential also exists for carbon sequestration through the land application of biosolids produced during VFA fermentation (73), which was beyond the scope of this study. As such, several long-term opportunities exist to advance VAF-SAF toward net-zero emissions as the technology develops with cost and carbon footprint targets in mind.

Conclusion
This work demonstrates the conversion of food waste–derived VFAs into SAF satisfying normal paraffins suitable for ASTM Fast Track qualification at 10 vol% blends, as well as highly branched isoparaffins produced via aldol condensation that can offer increased blending while maintaining ASTM fuel property specs. Statistical distribution models for VFA ketonization identified the VFA carbon chain length distributions suitable for each SAF upgrading pathway with stable ketonization catalyst time on stream performance observed for >100 h and minimal biogenic impurities in the final hydrocarbon fuel. Fuel property testing of the normal paraffin Fast Track VFA-SAF determined that flash point is the limiting fuel property, with a 20 vol% blend limit. Conversely, low-temperature viscosity was the limiting fuel property with Aldol Condensation VFA-SAF with 20 vol% blend limit. Due to their complementary fuel property limitations, synergistic blending was observed when combining Fast Track and Aldol Condensation of VFA-SAF to increase the overall blend limit to 70 vol%. Techno-economic analysis for VFA catalyst upgrading determined VFAs can be produced at $0.30/kg to achieve a Fast Track minimum fuel selling price of $2.50/gallon. Due to the potential for significant avoided methane emissions when diverting food waste from landfills for VFA production, life cycle analysis determined that VFA-SAF has potential to provide up to 165% reduction in greenhouse gas emissions relative to fossil jet fuel.

Materials and Methods
A brief description of the materials and methods is provided here, with additional details found in SI Appendix.

VFA Feedstock. VFA samples were provided by Earth Energy Renewables in their neat form following anaerobic digestion of food waste and recovery from fermentation media. Inorganic impurities were measured using inductively coupled plasma mass spectroscopy. Model VFAs for ketonization catalyst stability testing were obtained from Sigma-Aldrich.

Ketonization Kinetic Model. A simplified kinetic model was used to predict ketone profiles at 100% conversion of VFAs. Equations were solved using MATLAB software.

Catalytic Upgrading. VFAs were catalytically upgraded via ketonization over ZrO2 in a custom-built flow reactor that was outfitted with an online NDIR detector. Spinning band distillation was performed to separate ketones from fermentation media. Inorganic impurities were measured using inductively coupled plasma mass spectroscopy. Model VFAs for ketonization catalyst stability testing were obtained from Sigma-Aldrich.

Hydrodeoxygenation was performed in the custom-built flow reactor over Pt/Al2O3 for the neat ≥6 carbon ketone sample as well as the neat enone sample. Products were analyzed with a GC equipped with a Polyarc flame ionization detector and mass spectrometer.

Catalyst Characterization. ZrO2 catalyst used for VFA ketonization was analyzed by pyrolysis, NH3 temperature-programmed desorption, pyridine diffuse reflectance Fourier transform infrared spectroscopy, TGA, CHN, STEM-EDS, and XPS.

Fuel Property Analysis. Hydrocarbon analysis was performed by GCxGC, and distillation behavior was analyzed by GC. VFA-SAF fuel properties were estimated and measured using Tier α and β testing along with additional ASTM measurements conducted for the net heat of combustion, percent hydrogen, acid content, and nitrogen content. Sooting tendency was measured by yield sooting index.

Techno-Economic Analysis and Life Cycle Analysis. The process model was developed in Aspen Plus to determine the mass and energy balance. Model outputs were used for financial analysis. Catalyst costs were estimated using CatCost (68). Life cycle analysis was performed using data from GREET (74), SimaPro (75), and literature.

Data Availability. All study data are included in the article and/or SI Appendix.

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