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Elucidating the role of reactive nitrogen intermediates in heterocyclization during hydrothermal liquefaction of food waste

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Abstract:

Hydrothermal liquefaction (HTL) has emerged as a promising strategy for converting abundant, water-rich organic streams into an energy-dense, sustainable, biocrude. Despite major strides in improving oil yields and process intensification, a key underlying challenge persists in relating the composition of the feedstock to the ultimate fate of nitrogen, which is especially important for biocrude quality. To elucidate how nitrogen partitions into the respective gas, aqueous, oil and solid hydrothermal products, food waste was treated under HTL conditions (15 wt% solids, 575 K reaction temperature, and 1 hour reaction time) and elemental, chromatographic and mass spectroscopy analyses were performed on the products to establish dominant product classes and to close mass, carbon, and nitrogen balances. Liquid products (aqueous and biocrude) were found to contain nearly 60% of the nitrogen, with a majority of the biocrude-phase nitrogen in the form of amides. *Ab initio* thermodynamic and kinetic simulations on model reactants were used to evaluate potential reaction pathways involving reactive nitrogen intermediates. A subsequent kinetic model was evaluated at reaction temperatures ranging 300 – 1000 K and for a range of feedstock compositions to identify primary reaction pathways. The Maillard reaction is revealed to be the dominant pathway for converting reactive nitrogen intermediates into observed products, especially those in which nitrogen is present as a heterocyclic aromatic. Subsequent product analysis by Fourier Transform Ion Cyclotron Resonance Mass Spectrometry (FT-ICR MS) confirms strong agreement among the experimental data, kinetic model and proposed Maillard pathways.

Introduction

The current world reliance on fossil fuels and the corresponding environmental impacts have created global interest in alternative, sustainable energy sources. Particularly attractive are approaches that leverage renewable, distributed inputs to sustainably produce liquid fuels and chemicals. Hydrothermal liquefaction (HTL) has emerged as a promising candidate for converting water-rich mixed organic streams into biocrudes without extensive energy-intensive pretreatment. Processing water-rich feeds is typically challenging for thermal processes but is ideal for hydrothermal methods.¹ HTL's high level of feedstock flexibility^{2, 3} and the ability to directly produce a liquid biocrude has contributed to a recent surge of interest.³⁻⁵ HTL biocrude production has been demonstrated for algal, lignocellulosic, and waste feedstocks (**Table 1**) with energy recoveries ranging from 30 – 60%.⁶⁻⁸ Despite algae providing greater yields and energy recovery than other feeds, the high cultivation cost and ash content limit its effectiveness and have motivated research into HTL processing of biomass waste, sewage sludge, and food waste feeds.⁹⁻¹¹

Municipal waste streams (sewage sludge, food waste) are widely available, low-cost feeds that are rich in carbon.^{6,12, 13,10} Their water content precludes thermal conversion methods, other than hydrothermal ones. Use of waste feedstocks is economically and environmentally friendly due to low, potentially negative, cost and because the energy content of these feeds would otherwise not be used productively.^{12, 14, 15} Unfortunately, the heteroatom content of waste feeds, especially nitrogen, reduces biocrude quality and possibly yields.^{12, 14, 15}

Table 1. Biochemical composition and proximate analysis of common HTL feedstocks: algae, food waste, and sewage sludge.

Feedstock	Proximate Analysis (%)									Source
	Protein	Carb.	Lipid	Ash	Moisture	C (%)	H (%)	N (%)	O (%)	
Algae	7 - 63	9 - 54	13 - 55	4 - 22	65 - 80	50-62	7-10	1-10	17-35	Shakya et al. ¹⁶
Food Waste	4 - 22	40 - 70	6 - 24	0 - 6	61 - 86	40-60	6-10	1-6	25-55	Paritosh et al. ¹⁷
Sewage Sludge	2 - 42	25 - 52	0.1 - 41	2 - 34	70 - 90	35-50	5-9	3-8	33-60	Chen et al. ¹

To obtain a high-quality oil product, substantial efforts must be made to minimize oil-phase heteroatom (N, O, S) content. HTL input streams can contain anywhere between 1 and 10 wt% nitrogen for woody biomass and high-protein algae, and >50 wt% oxygen.^{18, 19} Oxygen is typically

found in the aqueous and gas phases in near equal amounts, whereas as much as 50% of the nitrogen present in the feed appears in the biocrude as a contaminant that must be removed by subsequent treatments.²⁰ Nitrogen removal is conventionally more difficult than oxygen removal, and removing nitrogen heterocycles is especially challenging.²¹⁻²³ Moreover, nitrogen interferes with catalytic acid sites intended for oxygen and sulfur removal, further complicating upgrading efforts.²⁴ Minimizing the nitrogen upstream in the feed or in the biocrude formed during hydrothermal processing is preferred over removing it during biocrude upgrading. In the hydrothermal step, it would be particularly advantageous to selectively control the fate of nitrogen—namely to be able to partition it into the gaseous, solid or aqueous phases, leaving the primary biocrude phase nitrogen-free. Our current knowledge of the chemical pathways that result in these respective fates, however, are substantially lacking with regard to the role of nitrogen in these reactive mixed feeds.

To better understand these pathways, thermodynamic and kinetic simulations are required. *Ab initio* calculations have been used on biopolymeric systems to understand key degradation steps during lignin depolymerization^{25, 26} and to reveal primary homolytic cleavage pathways during cellulose pyrolysis.²⁷ Notably, however, these studies are typically constrained to the primary reactions of single component, well-ordered feeds or surrogates that are not able to capture the more reactive secondary couplings of oligomeric, mixed component solvated environment that is prevalent during hydrothermal waste processing.

Current knowledge regarding HTL mechanisms are limited to lumped kinetic studies which rely on gas chromatography and mass spectrometry (GC-MS) analysis.^{8, 28-30} While this approach has proven valuable in providing insight into product phase distributions and general nitrogen pathways, it fails to describe the highly reactive condensed phase environment and particularly, the molecular-level intermediate pathways and barriers for driving nitrogen to each product phase.^{8, 31, 32} When considering food waste decomposition, it is typically considered to progress through two consecutive stages (Figure 1). In the first, the biopolymers (starches, proteins, fats) are hydrolytically decomposed into short-chained oligomers and monomers.¹¹ These small, solvated molecules are then free to couple in increasing degrees to form the aqueous (light), oil (heavy) and solid (char) products. With this perspective, the focus of the current study is not to understand the first decomposition stage. Instead, it is to better understand how the small

degradation products build back up to form the biocrude, and specifically how the nitrogen functionality, primarily from the proteins, becomes incorporated into the biocrude.

The composition of the reactive intermediate mixture is derived from detailed chemical analysis of the product fractions. The biocrude alone may contain thousands of unique compounds,³³⁻³⁵ causing analytical challenges, particularly with conventional GC-MS which provide molecular-level precision for volatile biocrude components but is not sufficient for the significant portion of the biocrude that is not volatile enough for GC analysis.³⁶ Fourier-Transform Ion Cyclotron Resonance Mass Spectrometry (FT-ICR MS) can provide molecular formula with isotopic accuracy on a molecular basis across the entire molecular weight regime of the product oil and is a promising technique for biocrude analysis.^{34, 35, 37} While FT-ICR MS has been used to advance scientific understanding of molecular pathways in pyrolysis of biomass,³⁸⁻⁴¹ its application to biocrude analysis – and especially to pathway discovery – is much more limited.^{34, 42} A combination of GC-MS and FT-ICR MS along with the known initial feed structures are used in this study to carefully select key functionality to be incorporated into chemical surrogates to be used in DFT studies to evaluate intermediate reaction pathways.

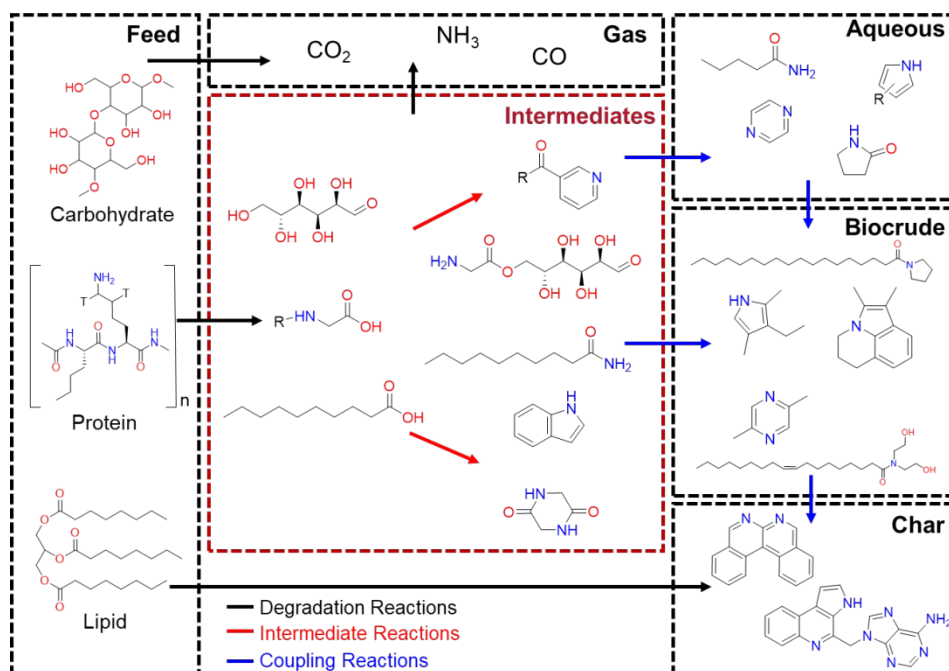


Figure 1. Representative nitrogen pathways for hydrothermally depolymerizing macromolecules of food waste and algae to reactive intermediates which form secondary condensation products before ultimately condensing to form biocrudes, and chars. This paper focuses on understanding the condensed phase chemistry in the reactions of intermediates (red dashed box).

This work will examine the role of feedstock composition and temperature on the type and amount of HTL product nitrogen. An analysis of small, volatile molecules found in food waste HTL biocrude will serve to identify important molecular classes which will be used to develop a reaction network for use in thermodynamic and kinetic simulations. The simulation results will then be compared to large biocrude molecules identified via FT-ICR MS to ensure model validity. The results of this work aim to assist in understanding nitrogen heterocycle formation at hydrothermal liquefaction conditions to make recommendations for optimal feedstock-specific operating conditions.

Materials & Methods

Development of Reaction Mechanism

Reaction mechanisms were formed capturing the interactions of the three primary components of non-woody biomass, i.e., protein, lipid, and carbohydrates. To maximize computational efficiency, simple model compounds were chosen. Glycine, the smallest amino acid, was chosen as a model protein degradation product; butadiene as the model fatty acid; and various small aldehydes (formaldehyde, glyoxal, furaldehyde, and glyceraldehyde) were used as model carbohydrate decomposition products. Glyceraldehyde was specifically chosen due to its known reactivity during the Maillard reaction.^{43, 44} HTL reaction mechanisms were identified from various experimental studies which involved identification of specific compounds resulting from macromolecular interactions.^{42, 45-49} Reactions were chosen to represent Maillard, hydrodenitrogenation and Aza Diels Alder (ADA) reactions to ensure representative understanding of the types of nitrogen reactions. These reactions were then divided into schemes based on the type and number of reactants involved.

Electronic Structure Calculations

Quantum chemical calculations were completed with the Gaussian 16 program suite. All chemical species are calculated in their singlet state and gas phase. The composite method CBS-QB3 was used to optimize geometries and evaluate single-point energies. Force constants and vibrational frequencies were calculated with B3LYP/CBSB7. The torsional motion around the single bonds in each molecule were treated as hindered internal rotations where each angle corresponding to an internal rotation was varied in incremental steps of 10° for 36 steps to achieve

complete rotation, allowing other coordinates to relax, evaluated with B3LYP/6-31G(d,p). Transition states were also calculated using CBS-QB3, with frequency calculations performed to verify the existence of only one imaginary frequency per transition state. When necessary, additional optimizations were performed to ensure the result was the lowest energy transition state. All calculations (optimization, frequency, and hindered rotor scans) were repeated using the density functional theory method M06-2X/6-31G+(d,p) for comparison but the results presented here are from CBS-QB3, apart from Scheme 4 reaction 3 because species M_{29} and transition state 4.3 could not be calculated at CBS-QB3 on the available computer hardware.

As water is the solvent and thus in extreme excess, it is taken to be pseudo zeroth order in all kinetic expressions. This is further confirmed by sensitivity analysis (**Figure S4**) which shows little sensitivity to changes in water concentration. Solvent effects can have significant impacts on some reaction rates, especially when reactants or products have strong interactions with the solvent, e.g. are hydrophobic or form hydrogen bonds.⁵⁰ For example, Bini et al.⁵¹ reported an increase in Diels Alder reaction rate in aqueous media, whereas Baisier et al.⁵² also found an increase in Maillard reaction rates in the presence of water with an excess of sugar, representing the two primary reaction types explored in this work. Furthermore, some reactions can be catalyzed by an explicit interaction with a solvent molecule. These effects are necessarily neglected in the current calculations, but the presumption is that to some extent the effects will alter all the competing pathways in a similar manner, and although individual energies and rates will be imprecise, to a first approximation the qualitative trends and relative predictions should be robust.

Kinetic & Thermodynamic Parameter Estimation

Enthalpies of formation are evaluated from the CBS-QB3 results using Bond Additivity Correction (BAC) values available in CanTherm Version 2.1.7.⁵³ To calculate partition functions, the CanTherm software package was utilized with a frequency scale factor of 0.99 for CBS-QB3 and 0.967 for M06-2X/6-31G+(d,p). The results from potential energy scans $V_i(\theta_i)$ are fit to a Fourier series in CanTherm and the resulting coefficients A_m and B_m in **Equation 1** are found using the least squares method.

$$V_i(\theta_i) = \sum_{m=1}^M A_{m,i} \cos(m\theta_i) + B_{m,i} \sin(m\theta_i) \quad (1)$$

Wherein V_i is potential energy change due to torsion i . $A_{m,i}$ and $B_{m,i}$ are fitting parameters and θ_i is the angle of rotation of the torsion i . The potential values were then used in solving the 1D Schrodinger equation to finally combine the harmonic oscillator (HO) approximation with the hindered rotor contribution. Entropy calculations considered molecular symmetry. Conventional transition state theory was used for calculating rate coefficients for considered pathways, with Eckart tunneling corrections to improve estimated rate coefficients at HTL-relevant temperatures. Rate constants were then fit to the modified Arrhenius equation (**Equation 2**) for each of the i^{th} reactions where k_i is the rate constant for each reaction, A_i is the pre-exponential factor, E_i is the activation energy, n is a modification factor to represent the changes more accurately with temperature, and T is the temperature in Kelvin.⁵⁴

$$k_i = A_i T^{n_i} \exp\left(\frac{-E_i}{RT}\right) \quad (2)$$

Kinetic Modeling

Thirty-eight reactions consisting of forty-seven molecules were proposed to describe initial intermediate nitrogen reactions in hydrothermal liquefaction. Arrhenius parameters for each reaction were calculated as described above then combined with their respective first and second order rate expressions to form a system of forty-eight ordinary differential equations (Equation 3). This system constituted the kinetic model, which was then integrated in MATLAB using ode15s given initial conditions corresponding to the fractional feed composition of the amino acid (M_1), lipid (M_7), and carbohydrate (M_{12} , M_{18} , M_{26} , or M_{35}). Kinetics calculations were completed as a function of temperature and reaction time. As discussed later, initial compositions covered the ternary compositional space defined in **Table 1**. Rates were calculated at 300-1000 K. All simulations were integrated over 30 minutes, unless denoted otherwise, to be consistent with typical experimental HTL timescales. Simulations did not consider secondary reactions to higher molecular weight polymers, chars or phase changes.

$$\frac{dc_j}{dt} = \sum_{i=1}^N \nu_{ij} \times r_i \quad (3)$$

$$r_i = k_i(T) \prod_{j=1}^M c_j^{-\nu_{ij}} \quad \text{for } \nu_{ij} < 0$$

Where c_j is the concentration of species j of $M = 47$ species; ν_{ij} is the stoichiometric coefficient of species j in reaction r_i of $N = 38$ reactions. Reaction rates were calculated with mass action kinetics based on the proposed mechanism using rate constants described in **Equation 2**, as summarized in **Table S3**.

Experimental Methods

Hydrothermal liquefaction experiments were conducted in a 300 mL stainless-steel Parr reactor (Model 4561) fit with a magnetic stirrer consistent with previous studies.^{9, 55} For each experiment, 100 g of food waste slurry (15 wt% solids) was loaded into the reactor. The food waste slurry was prepared as previously described^{9, 55} and according to Army specifications using a mixture of applesauce, chicken, butter, green beans, rice, instant potatoes, and cheese. After loading, the reactor was sealed and purged three times with nitrogen to remove residual air before pressurization to 65.5 ± 5 bar and heating to 575 ± 5 °C at approximately 279 K/min using an external heating jacket. After heating, the reactor pressure was 200 ± 5 bar, sufficient to maintain water in its liquid state. The reaction temperature was maintained for 60 minutes, at which point the reaction was quenched by placing the reactor in an ice bath until the measured temperature reached 310 ± 2 °C. Quenching required less than 10 minutes. The biocrude/solid mixture was then separated from the aqueous phase via vacuum filtration before the solids and biocrude were isolated using acetone. The acetone was then evaporated using a rotary evaporator to obtain the final mass of biocrude. Each product phase was carefully massed to ensure closure of the mass balance, as described previously.^{9, 55} Deliberate safety precautions were made to handle the high pressure and high temperature reaction mixture. All experiments were conducted in a ventilated fume hood with a properly sized rupture disk (172 bar) vented to the hood. The reactor was pressure tested with nitrogen at ambient temperature to 69 bar prior to each run.

All runs were completed in triplicate, with yield measurements agreeing to within $\pm 10\%$ when experiments were performed under identical conditions. Average values obtained from these experiments are presented here.

HTL biocrude was analyzed via GC-MS, to understand the molecular composition of the volatile fraction. Biocrude and aqueous samples were dissolved in toluene to a final concentration of 125 $\mu\text{g}/\text{mL}$ for positive-ion APPI Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR MS). Dissolved samples were analyzed using a custom-built FT-ICR mass spectrometer⁵⁶, equipped with a 22 cm room temperature bore 9.4 T superconducting solenoid magnet (Oxford Instruments, Abingdon, U.K.) and a modular ICR data collection station (Predator).³³ Both techniques are described further in the **Supporting Information**. Further analysis was completed to close the mass balance for carbon, nitrogen, and oxygen. Total organic carbon (TOC) and total nitrogen (TN) analyses were completed on the aqueous phase using a TOC analyzer (Shimadzu Co. Kyoto, Japan). Elemental analysis of the biocrude and char phases was completed by Midwest Microlabs (Indianapolis, IN). The hydrogen and oxygen composition of the aqueous phase was taken to be the difference between the total feed oxygen content and the sum of the remaining products.

Results and Discussion

In this work, reaction pathways for the liquefaction of carbohydrate, protein, and lipid monomers are explored through an analysis of thermodynamic reaction equilibrium and kinetics. GC-MS analysis of biocrude from food waste HTL is used to gain an understanding of small biocrude-phase products. A density functional theory (DFT) model is used on proposed reaction schemes involving six primary pathways for the formation of HTL intermediates. Free energy and Arrhenius parameters are calculated and utilized in the development of a kinetic model to evaluate the rates at which these reactions occur. This analysis provides a baseline for understanding the feasibility of the proposed pathways via a discussion of the coupling reactions from the primary model products to the larger molecules seen in HTL biocrude. FT-ICR MS was employed to validate model predictions through comparisons between the carbon number and double bond equivalency of computationally-predicted and experimentally-observed product molecules.

Experimental Analysis of Volatile HTL Products

To ascertain the role and molecular contributions of nitrogen during the HTL process, a food waste slurry consisting of 18% protein, 59% carbohydrate, and 22% lipid was heated to 575 K for 60 minutes at 200 bar. This reaction produced, on average, 14% biocrude, 25% solid, and

17% aqueous products on a mass basis. Resultant mass yields were used in conjunction with the elemental composition of each phase to close carbon, nitrogen, hydrogen, and oxygen mass balances within 10%. This analysis shows that the feed contains 3.8% nitrogen, of which, 13.2% is found in the biocrude. The nitrogen product fractionation for food waste HTL can be seen in **Figure 2**, wherein nearly 50% of the nitrogen partitions to the aqueous phase, consistent with previous literature findings.^{20, 57} The gas phase was found to contain >98% CO₂, consistent with prior work showing only trace amounts of gas-phase ammonia.^{58, 59} Gas-phase nitrogen is below detection limits, leading to it being lumped together with loss, resulting in 95% mass balance closure. Nearly 60% of the nitrogen can be found in the liquid products (biocrude and aqueous), which are assumed to be miscible at reaction conditions, but phase partitioned at ambient temperature. These two phases have the potential to undergo the same reactions before thermodynamic partitioning occurs upon quenching.

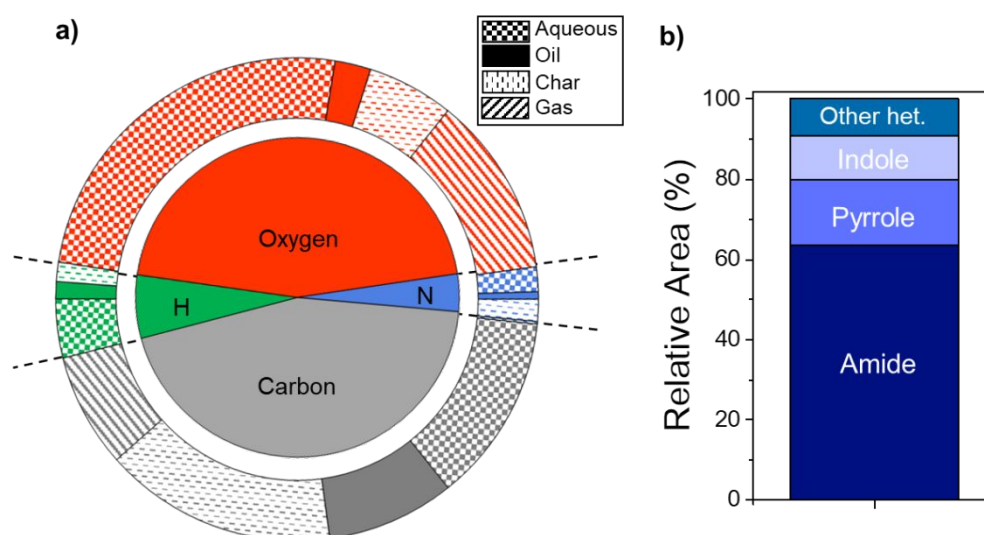


Figure 2. a) Elemental analysis tracking the fate of carbon, hydrogen, oxygen, and nitrogen from food waste feedstock (inner circle) to HTL products (outer ring). HTL was performed at 300 °C, 1 h residence time. **b)** GC-MS peak area percentages for the total nitrogen identified in the oil phase, broken into prominent classes. Indoles, pyrroles, and other N heterocycles can be grouped together as the total nitrogen heterocycle content.

Further analysis utilizing GC-MS (**Figure S1, Table S1**) of the volatile portion of biocrude allows for molecular-level analysis of small molecules where 86% of the total peak area was identified as fatty acids. Fatty acids were confirmed as the most abundant product via GC×GC-HRT, further identifying a suite of fatty acid amides corresponding to the same range of carbon chain lengths (**Figure S2**). The second most abundant class is nitrogen-containing compounds, of which 64% have been identified as amides (**Figure 2b**). Further analysis of the chromatogram

identifies a range of pyrroles, pyrrolidines, and indoles, all classes of nitrogen heterocycles, which accounted for the remaining 36% of the nitrogen compounds identified. Pyrroles account for nearly half of the identified heterocycles and indoles for an additional 30%. These compounds contain nitrogen locked in a ring structure, making them difficult to upgrade and remove. Their presence in HTL biocrude is of express interest in this work.

Simulation of Reaction Intermediates

Due to the short-lived nature of the reactive small molecules and computational limitations, model species representative of the protein, carbohydrates, and lipid degradation products from food waste were chosen as feedstock molecules. Additional carbon atoms will serve to increase the hydrophobicity at room temperature and can play a role in increased steric hinderance that can affect experimental reaction rates. For this study, glycine was utilized as the model protein, butadiene as the model fatty acid, and four aldehydes (formaldehyde, glyoxal, furaldehyde, and glyceraldehyde) as model carbohydrates. Reaction chemistries commonly identified in literature were evaluated and compared including combination (condensation, cyclization, dehydration) and decomposition (hydrolysis, deamination, decarboxylation) reactions.

At HTL conditions, inorganic and small organic nitrogen molecules readily break down into amines at high temperatures in basic media through various deamination pathways.⁶⁰ Heterocycles, such as pyrroles and indoles, which are present in the side chain of amino acids proline, histidine, and tryptophan, lock their nitrogen in a stable aromatic ring structure which requires severe reaction conditions to break down. The Maillard reaction is a well-known heterocyclic reaction responsible for the browning of foods.^{61, 62} Maillard reactions occur between a carbonyl group on a reducing sugar and the amine group of a protein, and have been shown to proceed through pH-dependent pathways.^{63, 64} Specifically, retro-aldol reactions are more prominent at basic pH, producing more reactive C₂ – C₅ sugar fragments.⁶³ An additional reaction of importance is the Aza-Diels Alder (ADA) reaction, which has not received as much research attention as the well-known Diels-Alder reaction. ADA represents the cycloaddition of a diene and dienophile without an imine group.^{65, 66} This reaction is thought to occur via a single transition state without intermediates. Based on HTL literature and established chemical pathways, the final reaction scheme was reduced to those shown in **Figure 3**.^{20, 31, 42, 43, 65, 67-75} Studied reactions are divided

into six primary pathways denoted by their initial reactants, and all molecules are summarized in **Table S2**.

Scheme 1 is the result of glycine decomposition in four potential competitive pathways including decarboxylation (1.5) and deamination (1.3). Schemes 2 – 4 involve the addition of increasingly oxygenated aldehydes (formaldehyde, furaldehyde, glyceraldehyde). Each of the aforementioned schemes include a pathway to the primary amine and a competitive reaction (2.3, 3.3, 4.3) resulting in nitrogen heterocycle formation through the ADA reaction mechanism. The remaining secondary reactions in these schemes do not result in heterocycle formation but include rearrangements, aminations, dehydration, and decomposition reactions. Scheme 5 represents a modified Maillard reaction for the formation of a nitrogen heterocycle. This is an important reaction to consider due to its prevalence in literature and as a dominant heterocycle formation mechanism. The Maillard reaction contains ten intermediates, making it the longest scheme with multiple kinetic barriers and free energy increases to overcome. The last scheme considered, (6) outlines the pathway for the combination of two glycine molecules through dehydration to form a nitrogen heterocycle.

Transition state theory is used to calculate the temperature-dependent reaction rates while accounting for tunneling effects and fit to a modified Arrhenius equation (**Equation 2**) as reported in **Table 2**. The Eyring equation is then used to derive the Gibbs energy of activation ($\Delta^\ddagger G^\circ$) at 600 K, as described in the Supporting Information. This strategy allows for the evaluation of the free energy change for the reaction, indicating the favorability of the reaction, as well the activation free energy, representing the ΔG of the transition state. An exergonic decrease in free energy signifies the system releases energy spontaneously and is favorable in comparison to an endergonic increase in free energy where further energy inputs are required. Likewise, a high activation free energy signifies a slow reaction, and a low activation free energy signifies a faster reaction. Transition state activation free energy and free energy of reactants are shown on the free energy surfaces in **Figure 4** and tabulated in **Table S3**, all evaluated at 600 K. Reaction schemes are colored based on the reaction step with the largest activation energy, except when the largest step is a rearrangement reaction. The $\Delta^\ddagger G^\circ$ values at 600 K are also shown in parentheses next to the reaction numbers in **Figure 3** and are the activation barriers described in the following discussion.

In Scheme 1, the deamination and decarboxylation of glycine in reactions 1.4 and 1.5 are the most exergonic first-step reactions, however, reaction 1.4 has the highest activation barrier of the reaction network at 109 kcal/mol. The decarboxylation in reaction 1.5 has the lowest barrier of the initial competitive steps in Scheme 1 at 74 kcal/mol. Reaction 1.1 and 1.3 also boast high activation barriers, both at 107 kcal/mol. Glycine decomposition through the deamination pathway (1.4), despite its activation energy, has been shown to occur at temperatures as low as 559 K, making this pathway the most probable first reaction step in Scheme 1.⁷⁶

In Scheme 2, the formation of a nitrogen heterocycle through ADA is kinetically limited by reaction 2.2 in which the hemiaminal formed in the previous reaction undergoes dehydration to a Schiff base. This reversible reaction becomes increasingly exergonic (and hence spontaneous) as temperatures rise above about 400 K but has the highest barrier in Scheme 2 (57 kcal/mol) and a rate strongly dependent on reaction temperature. The ADA reaction (2.3) occurs with a slightly lower energy barrier (52 kcal/mol) than the competing reaction. Reaction 2.3 also has a lower energy barrier than the competing decarboxylation reaction, indicating heterocycle favorability. The same analysis holds true for Schemes 3 and 4, wherein ADA has a lower activation barrier than the competing reactions studied. The predicted nitrogen heterocycles from ADA are all substituted pyridines, wherein GC-MS analysis does not identify any pyridines but instead has a higher concentration of nitrogenated five-membered rings. This may be due to the presence of tryptophan in real food waste that is not considered in the model.

In Scheme 5, many of the reactions have energy barriers above 50 kcal/mol. Reaction 5.2 of the Maillard reaction has a large activation energy at 61 kcal/mol and reaction 5.5 has the largest energy barrier at 66 kcal/mol. Reaction 5.2, the formation of a Schiff base, is reversible, but at typical HTL conditions the reverse rate is negligible, especially compared to the onward reaction 5.3, and has not been considered. The Strecker Degradation (SD) that occurs in the final stages of the pathway (5.2) as well as the dicarbonyl produced from the Amadori rearrangement (5.4) react with an amino acid (glycine) to produce a hemiaminal (5.7) via a mechanism with an energy barrier of 54 kcal/mol, after the second rate-limiting step at 5.5. This reaction is the dehydration of a previously formed nitrogen heterocycle (1,2,3,6-tetrahydropyrazin-2-ol) to 2,5-dihydropyrazine. Reaction 5.8 is necessary for nitrogen heterocycle formation, followed by two dehydration rearrangements. Due to the large energy barrier of reaction 5.9, it is most likely that piperazine-

2,5-diol (M_{43}) will be the final product of this reaction scheme. Experimentally, only one pyrazine compound was identified with GC-MS constituting 2% of the nitrogen-containing compounds. Contrary to the model predicting alcohol-substituted pyrazine, the observed molecule is methylated, potentially indicating the prevalence of tertiary dehydration reactions at longer residence times. On the other hand, the coupling of two glycine molecules in Scheme 6 via dehydration occurs with an energy barrier of 64 kcal/mol, indicating the enhanced stability of glycyglycine (M_{46}) and the potentially unfavorable dehydration to piperazine-2,5-dione (M_{47}).

Another consideration, across all schemes, is the formation of gaseous byproducts. CO_2 is produced in reactions 1.5, 2.4, 3.4, and 5.3 while NH_3 can be found in reactions 1.3 and 1.4. All CO_2 formation reactions considered are thermodynamically favorable at 600 K, constituting a decrease in free energy, whereas only one of the two ammonia producing reactions is exergonic. Kinetically, all reactions forming CO_2 as a byproduct have lower activation energies than the two NH_3 reactions by at least 33 kcal/mol. This finding is consistent with experimental results, where over 98% of the gaseous products are carbon dioxide, and only trace ammonia is detected.

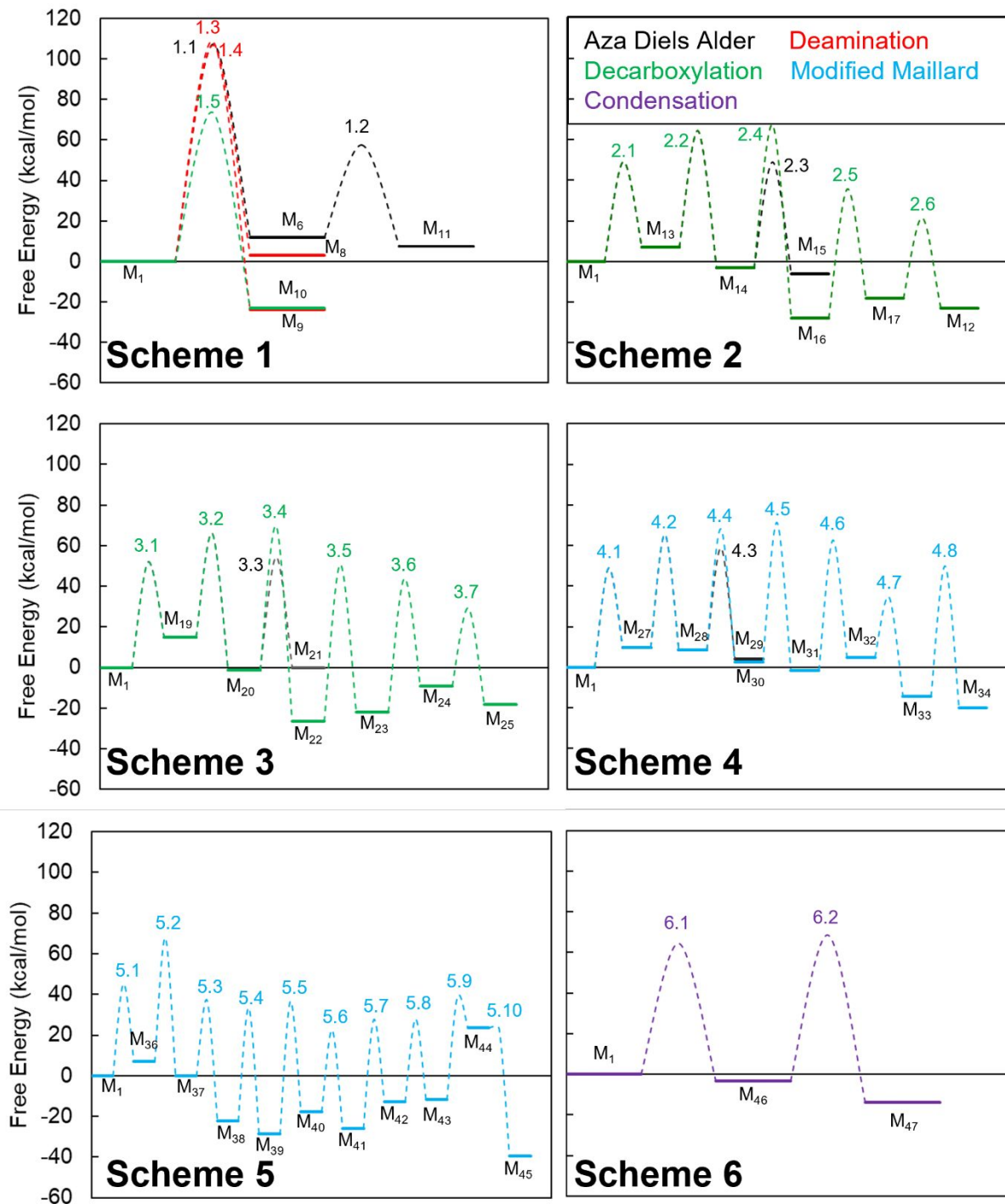


Figure 4. Free energy diagram for all reaction pathways at 600 K. Reaction pathways are colored based on the reaction step with the largest activation energy, except when a rearrangement reaction had the highest activation energy.

Table 2. Modified Arrhenius rate parameters (Equation 2) calculated using the CBS-QB3 method. Calculated rate constant for all reactions at 600 K.

Reaction	A_i	n	E_i (kcal/mol)	k_i (600 K)	Class ⁺
1.1*	2.4 E-79	27.34	64.1	1.0E-26	Degradation
1.2**	1.8 E+04	1.53	20.3	1.3E+01	ADA ⁺
1.3**	4.0 E+01	3.04	85.7	6.6E-22	Deamination ⁺
1.4**	4.7 E+06	2.07	94.1	1.3E-22	Deamination ⁺
1.5*	1.1 E-24	11.07	56.2	2.0E-14	Decarboxylation ⁺
2.1**	4.6 E-05	4.29	21.2	7.1E-01	Combination
2.2*	1.0 E-06	5.71	48.6	1.5E-08	Dehydration
2.3**	3.0 E-01	2.69	22.6	5.3E-02	ADA ⁺
2.4*	4.6 E-33	13.58	49.3	2.6E-13	Decarboxylation ⁺
2.5**	6.2 E+01	3.12	43.6	3.8E-06	H ₂ O Addition
2.6*	5.9 E+02	2.86	33.2	4.4E-02	Degradation
3.1**	1.3 E-06	4.65	22.4	7.6E-02	Combination
3.2*	3.3 E+18	-1.57	54.0	3.1E-06	Dehydration
3.3**	5.7 E-01	2.32	23.9	3.0E-03	ADA ⁺
3.4*	1.3 E-31	13.03	49.9	1.3E-13	Decarboxylation ⁺
3.5*	2.9 E-07	5.86	68.3	7.4E-16	Rearrangement
3.6**	3.8 E-01	3.48	42.3	6.9E-07	H ₂ O Addition
3.7*	4.6 E+12	-0.06	36.8	1.3E-01	Remove CH ₂ O
4.1**	1.5 E+01	2.21	20.2	9.0E-01	Combination
4.2*	8.0 E+08	0.91	51.2	6.2E-08	Dehydration
4.3**	4.5 E+01	-0.67	25.5	3.3E-01	ADA ⁺
4.4*	5.7 E-14	8.42	51.4	2.6E-09	Rearrangement
4.5*	7.6 E-01	3.43	58.9	9.3E-13	Rearrangement
4.6*	3.2 E-30	12.83	45.1	5.3E-11	Rearrangement
4.7*	1.8 E+16	-1.13	29.7	2.0E+02	Degradation
4.8*	5.5 E-47	18.11	39.5	4.6E-11	Rearrangement
5.1**	7.7 E-04	3.75	16.6	1.8E+01	Combination
5.2*	9.3 E+17	-1.31	64.6	6.5E-10	Dehydration
5.3*	1.4 E+12	-0.25	32.9	2.9E-01	Decarboxylation
5.4*	1.4 E-29	12.66	37.0	6.7E-08	H ₂ O Addition
5.5**	4.0 E-20	9.16	33.4	7.9E-07	Decomposition
5.6*	3.6 E+13	-0.24	39.7	2.7E-02	Addition
5.7**	5.7 E-07	4.44	21.6	1.7E-02	Rearrangement
5.8*	8.3 E+09	-0.15	31.0	1.7E-02	Dehydration
5.9*	2.5 E+15	-0.35	54.2	4.6E-06	Dehydration
5.10*	1.1 E+13	0.44	3.0	1.5E+13	Dehydration
6.1**	2.0 E-01	2.87	35.5	2.2E-06	Coupling
6.2*	1.7 E+06	1.23	62.3	8.9E-14	Condensation ⁺

*1st order reaction with units A [=] s⁻¹, k [=] s⁻¹**2nd order reaction with units A [=] cm³/mol*s, k [=] cm³/mol*s⁺ Indicates primary reaction mechanism corresponding to colors in Figure 4.

The kinetic parameters summarized in **Table 2** are used to construct a model to perform a reaction flux analysis and identify dominant chemical pathways during hydrothermal operation, particularly with the goal of revealing pathways by which the observed heterocyclic nitrogen molecules are produced. The model does not consider further polymerization reactions, solely predicting the behavior of the intermediates. Cross reactions, wherein products from one scheme can react with reactants or products from another scheme, were also not considered.

The molecules considered in this reaction network are distributed into fourteen chemical classes (**Table S2**) for ease of reporting the kinetic model. The first three classes denote the primary reactants: amino acid, diene, and aldehyde which together account for six of the forty-seven molecules examined. Additional chemical classes identified are as follows: Amine, Amide, Hemiaminals, Amino-aldehydes, Alkanolamines, Acids, Schiff base, Gases, Nitrogen heterocycles, and Other. These classes will be referenced throughout the remainder of this study.

Temperature Dependence

Standard HTL reactions take place between 525 – 675 K, resulting in a range of potential dominant reactions. Reaction temperature has been shown to affect HTL mass yields and composition.³⁴ One variable often considered is the optimal reaction temperature. Operating below 525 K often results in high solids (char) formation and is traditionally coined hydrothermal carbonization. The model assumes and kinetics confirm that the monomeric species are stable at these reaction temperatures, and will not couple to form oligomers, biocrude-soluble products, or char. This observation is consistent with the understanding that carbonization primarily progresses directly from concerted biopolymer rearrangements, not from the bottom-up approach whereby the biocrude is polymerized to char.⁷⁷ Hydrothermal liquefaction reactions, on the other hand, typically occur between 525 – 625 K to ensure complete breakdown of macromolecules and to limit low-temperature char formation. With this analysis, operating conditions can be fine-tuned to limit heterocycle production in the liquid products. **Figure 5** plots the effect of reaction temperature at 30 minutes for four real feedstock compositions taken from algae and food waste (**Table S4**).^{9, 16} Carbohydrate concentration was divided evenly among the four possible reactant aldehydes. All kinetic simulations were assumed to be single phase in this study, and changes to

physical or kinetic properties upon phase transition (oil/aqueous or supercritical) were not considered.

Low protein feeds (< 50%) resulted in hemiaminals as the sole nitrogen intermediate product at HTL conditions (525 – 625K). At these conditions, higher protein feeds also contain a small (< 3%) fraction of amides that increases substantially with increasing temperature. When the majority of the feed is carbohydrates, no amide formation occurs at any of the explored temperatures, potentially due to the low concentration of glycine, inhibiting reaction 6.1 due to its high kinetic barrier compared to parallel steps in the competing schemes. There is no shortage of aldehydes to undergo Maillard and competing combination reactions. The model predicts an overwhelming majority of hemiaminals within traditional HTL temperature range, yet a small increase in temperature introduces unfavorable heterocycles as well as amide, gaseous, and amine products.

Despite predicting no appreciable formation of nitrogen heterocycles until 650 K, experimental findings from algae and food waste feeds have identified oil-phase heterocycles at 575 K. This is likely caused by a combination of unreactive side chains present in the native proteins and tertiary reactions whereby the intermediate products continue to react as they build up to larger biocrude molecules. Five of the twenty essential amino acids contain heterocyclic side chains and are likely present in food waste. In experiments performed between 550 – 590 K with algae, appreciable amounts of pyrroles, pyridines, indoles, diketopiperazines (DKPs), and imidazoles were identified in the biocrude.^{34, 78} Previously mentioned analysis with food waste HTL has also identified the presence of nitrogen heterocyclic compounds, despite reaction temperatures below 650 K.³ The hemiaminals predicted by the model exhibit similar functionality to some of the observed heterocycles, including DKPs with the presence of nitrogen and double-bonded oxygen in the same molecule. Both have the potential to form from the interaction between amino acids and monosaccharides, often proceeding through a hemiaminal intermediate before the final cyclization step.⁷⁹ The experimentally observed heterocycles are larger (>C₆) and likely result from secondary coupling reactions from the intermediate products predicted here. Additionally, the model predicts less than 2% of the products are heterocycles in the typical reaction temperature range. Notably, the vast majority of the nitrogen remains as unreacted glycine due to the initial energy barriers to decomposition. The proposed model serves to understand dominant pathways

and molecular functionality that are critical steppingstones towards further combination and cyclization interactions to produce the experimentally identified molecules.

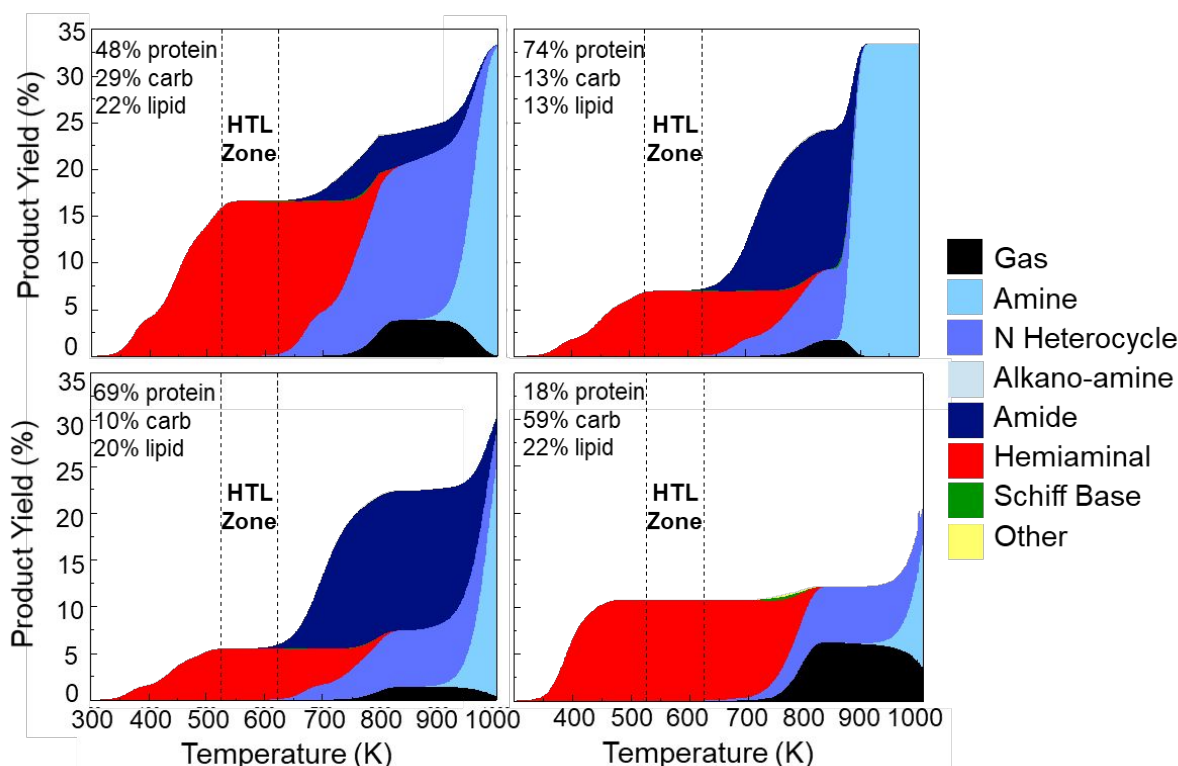


Figure 5. The effect of temperature on HTL intermediate product distributions for four surrogate feedstock compositions representative of food waste, and various algae streams. Shades of blue represent nitrogen containing classes identified in the GC-MS. Dashed lines indicate the temperature range for conventional HTL experiments. All models were completed at 30 min residence time for a range of 300 – 1000 K discretized to 1200 points.

Feed Composition Effects

The composition of the feed can vary from 2 – 75% protein (**Table 1**), which can have a substantial effect on the prevalence of nitrogen reactions and resultant product distributions. Due to this large variance, **Figure 6** examines the effect of feedstock composition on the concentration of four nitrogen-containing product classes (amine, hemiaminal, amide, and nitrogen heterocycles). The kinetic model was run at 600 K for 30 minutes with starting feed compositions evenly distributed between the reactants in 1% discretization. Due to the exclusion of secondary coupling reactions of intermediate products, concentrations do not continue to change (within 1%) after this time for all simulations. Product classes were grouped so as to exclude the initial

concentrations of amino acid, aldehyde, and diene. The resulting data was interpolated and presented as a heat map for the four heteroatom classes.

The amino acid, specifically glycine, is the sole reactant containing nitrogen in the model, leading to its concentration having the greatest effect on product distribution and yield. In the amine and amide classes, this trend is the clearest, where increased amino acid content directly increases resultant product concentration. The hemiaminal and nitrogen heterocycle classes, however, result in a more interesting trend that does not directly correlate with increasing amino acid concentration, indicating these molecules are the result of emergent monomeric interactions between macromolecule decomposition products. The hemiaminals (carbinolamines) are characterized by a hydroxyl and amine group attached to the same carbon, thereby opening additional pathways to formation including from aldehydes. Looking back to the reaction network, hemiaminals are products of the first reaction in Schemes 2, 3, and 4 (reactions 2.1, 3.1, and 4.1), which represents the addition of glycine with each of the starting aldehydes. It is for this reason that the highest concentration of hemiaminals occurs close to a 50:50 mixture of amino acid and aldehyde, continuing to decrease moving further away from the stoichiometric ratio.

While hemiaminal formation is relatively easy to explain, nitrogen heterocycle synthesis is more complicated. In the GC-MS analysis, HTL of food waste resulted in 36% of the volatile nitrogen species in the biocrude as heterocycles. Nitrogen heterocycles are formed in the reaction network in all schemes through a variety of pathways. Scheme 1 uses the Aza Diels Alder reaction to complete a cycloaddition of an imine (M_6), formed from the dehydrogenation of the amino acid, with butadiene (M_7). Schemes 2 – 4, however, react through the previously discussed synthesis of a hemiaminal (M_{13} , M_{19} , M_{27}) which then undergoes dehydration to produce the reactive imine (M_{14} , M_{20} , M_{28}). The Maillard reaction is responsible for nitrogen heterocycle formation in Scheme 5, wherein glycine undergoes an addition reaction with glyoxal (M_{35} , aldehyde) to form a hemiaminal (M_{36}) before dehydration to produce an imine (M_{37}). In Schemes 2 – 4, the same first three steps occur, with the addition of diene to promote nitrogen heterocycle formation. Scheme 5, on the other hand, does not include the addition of butadiene and instead allows for decarboxylation of the imine and subsequent hydrolysis and decomposition reactions leading to the formation of a Strecker aldehyde (M_{12}) and a reductive amination product. In the last stage, condensation of the two amino-carbonyl compounds results in the formation of a nitrogen

heterocycle (piperazinediol, M_{43}) that can further dehydrate to pyrazine compounds (M_{44} , M_{45}). Lastly, Scheme 6 produces a nitrogen heterocycle through two subsequent dehydration and inter/intramolecular condensation reactions to form a diketopiperazine (M_{46}).

In the kinetic analysis, Scheme 1 is not predicted to contribute substantially to the observed product distribution or heterocycle formation, likely due to the high activation barrier of reaction steps 1.1, 1.3, and 1.4 coupled with their slow rate constants on the order of 10^{-50} s^{-1} or $\text{cm}^3/\text{mol}^* \text{ s}$. The rate constants of schemes 2-6 are comparable, indicating their similar reactivity and likelihood for these reactions to proceed. For this reason, Scheme 5 appears to contribute the most to heterocycle formation due to its sustained low activation barriers, assuming the energy input is suitable to proceed past reaction 5.2 ($\Delta^\ddagger G^\circ = 61.1 \text{ kcal/mol}$).

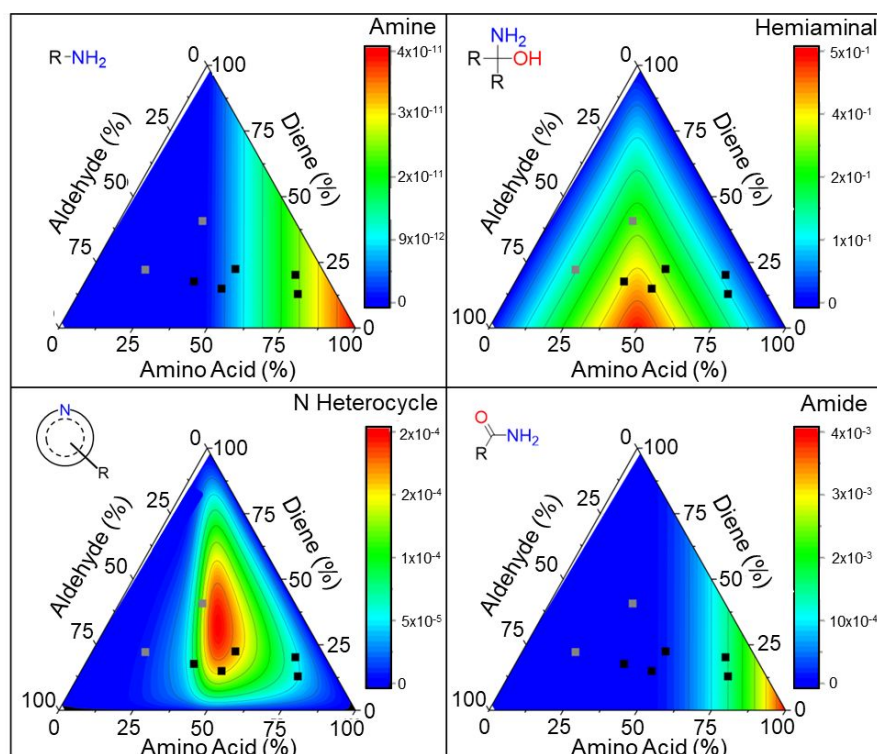


Figure 6. Ternary diagrams relating the product composition (color map) of four intermediate compound classes resulting from a particular starting composition defined on the ternary diagram. All data were taken interpolated from the kinetic model simulations at 600 K and 30-minute residence time. Black squares (□) represent known algae feedstock compositions and gray squares (□) are representative of food waste feedstocks where the aldehyde, amino acid and diene fractions are taken as surrogates for the carbohydrate, protein and fats, respectively. Color bar represents product concentration resulting from the kinetic model, [mol/cm³].

To further understand the effect of feedstock composition on nitrogen heterocycle formation, real feed compositions are referenced. In **Figure 6**, the black squares are representative

of various algae feedstocks while gray show food waste compositions, all highlighted in **Table 1** and **S3**. The comparatively low concentration of amines is primarily due to their role in the reaction scheme, wherein it is present as the reactant or as only one of the products of reaction 2.6. The lack of observed amines is indicative of their highly reactive nature in the reaction network and is consistent with GC-MS results which identified no primary amines in the product mixture (**Figure 2b**).

Heterocycles are the sole class, of the four examined here, whose concentration depends on all three primary macromolecule contents. For this reason, the center of the high intensity area appears to occur near a stoichiometric mixture of all three reactants. All five algae compositions are within the bounds of increased heterocycle production. Food waste, on the other hand, often contains low protein and high carbohydrate content, indicative of the typical American diet. Of the two food waste feedstocks explored, one occurs in a region of moderate protein concentration while the other is in an area of low protein concentration. Recall, heterocycles are highly undesired in biocrude, as it requires harsh hydrodenitrogenation (HDN) conditions to remove this nitrogen during upgrading. Elevated aromatic nitrogen has generally been observed in HTL oils from algal feeds, but lower amounts from food waste.^{80, 81} This is consistent with our model findings which show that sub-stoichiometric mixtures of protein results in lower heterocycle concentration. Cheng et al. has shown through FT-ICR MS that nitrogen heterocycles are present in both high and low-lipid algae biocrude in similar concentrations.³⁴ *N. salina*, a high-lipid algae, has shown to produce an oil high in alkyl substituted pyrroles and indoles, whereas *G. sulphuraria* biocrudes show higher degrees of alkylation with larger double-bond equivalency (DBE) range.³⁴ Low-lipid food waste, on the other hand, results in indoles as the primary heterocycle- still with an overall concentration less than 1%. This is most likely due to the presence of tryptophan as a common amino acid in chicken. Overall, the feed composition can be shown to be an important factor in determining the HTL product distribution.

Closing the Gap between Small and Large Molecules

Making the connection back to experimental results, **Figure 7** showcases the observed trends between all nitrogen-containing heteroatom classes for both the biocrude and aqueous phases. Positive ion APPI FT-ICR MS was utilized to probe the molecular space, revealing similarities in the carbon number and DBE relationship for both liquid phases (biocrude and

aqueous) across heteroatom classes. DBE provides insight into chemical makeup by capturing the degree of unsaturation resulting from double bonds or rings, as is typically resultant from molecular condensations. All four nitrogen classes contain biocrude molecules with a DBE greater than five, indicating the presence of at least one ring structure. It can also be seen that with increasing nitrogen number, the lower DBE limit increases, indicating further formation of heterocycles, most likely via dimerization pathways. Tryptophan, with an indole side chain, has the potential to undergo the Maillard reaction to produce a molecule with two nitrogen and a DBE > 10, consistent with identified molecules in the N_2O_x heteroatom class. Dimerization of this molecule then results in a compound with DBE > 20, $>C_{30}$ and four nitrogen, also consistent with **Figure 7**. Notably, over 70% of all observed nitrogen-containing compounds had $<C_{40}$, suggesting that chain growth may be limited based on the availability of reactive amines and molecules larger than C_{50} begin to partition to solid products.

Further comparisons of this data to macromolecular trends can be found in **Figure S3**, wherein the regions of high abundance are consistent with the polymerization of common carbohydrates and fatty acids. A compelling trend emerges from this data, whereby a strong linear relationship is observed across product classes. To better understand this trend, a range of chemistries were analyzed, starting with the Maillard reaction. By considering the chemistry that occurs during the Maillard reaction, it is noted that an amine condenses with a sugar molecule to form an N-substituted glycosylamine which undergoes Amadori rearrangement (5.4) to produce an Amadori rearrangement product (ARP). This is followed by a dehydration to produce furfurals and reductones, based on pH. Additionally, sugar fragmentation and Strecker degradation occur nearly simultaneously alongside the dehydration. During Strecker degradation, amino acids are oxidized in the presence of dicarbonyl compounds to produce CO_2 , aldehydes, and amino-ketones. The resultant aldehydes react with each other through aldol condensation in the final stage of the Maillard reaction. Finally, aldehydes react with amines through condensation and cyclization reactions to produce nitrogen heterocycles, including pyrroles, pyridines, and pyrazines. The Maillard reaction nets the loss of a carbon dioxide and three water molecules, with the addition of three $R-NH_2$ groups, effectively increasing the DBE by one (ring formation) and increasing the net carbon number by two, on average ($DBE = 0.5 \times C$). To better approximate this trend, chemistries from literature were examined^{34, 49, 82} by quantifying the change in DBE upon completion of the Maillard reaction as $[(DBE)_{products} - (DBE)_{reactants}]/[(C)_{products} - (C)_{reactants}]$; after doing this, the

average relationship was updated as $DBE = 0.48 \times C$, as plotted in **Figure 7** as a black dashed line. Previous work has identified the importance of the Maillard reaction in biocrude formation from model sugars and amino acids, citing the formation of pyrazines as well as increased production of aromatic amines and amides.⁴⁹

A similar analysis was performed using the reaction chemistry examined in this work. Reaction pathways were weighted by product abundance and correlated to DBE to reveal a trend, $DBE = 0.51 \times C$, with the corresponding fit plotted in green in **Figure 7**. For comparison with experimentally observed products, the relationship is extrapolated to larger carbon numbers as the expectation is that the intermediates will couple to form the larger products in secondary reactions. Notably, this relationship holds true as the dominant trend from all three data sets: the monomeric reactions (kinetic model), the final aqueous, and the final oil products, suggesting that the Maillard reactions are a dominant coupling pathway. Furthermore, the trend underscores that for every two carbons, one DBE is introduced, suggesting that other pathways may also be involved, such as aldol condensation or cycloaddition. As can be seen in **Figure 7**, the resultant model products correlate well with Maillard reaction products, consistent with the high degree of aromaticity observed in GC-MS. The trend also agrees with the identified pathways from the proposed intermediates to the highly aromatic compounds identified, indicative of aqueous phase molecules having a lower carbon number and DBE than oil molecules, with considerable overlap in their ranges. This suggests a common pathway for the formation of biocrude and water-soluble molecules, and is consistent with the understanding that at reaction temperatures, the two phases are miscible and are only phase separated and partitioned upon cooling.⁹ The FT-ICR MS findings show that polymerization is scale-independent where the ring structure and intra-chain unsaturations are maintained across small molecules in the aqueous and larger organics in the oil. Furthermore, the trend suggests a high degree of saturation, as in the monomeric species glycine, tetrahydropyridine, and N-methylmethanimine captured by this kinetic model. This serves as valuable insight into the validity of the model in accurately predicting key HTL intermediate product distributions.

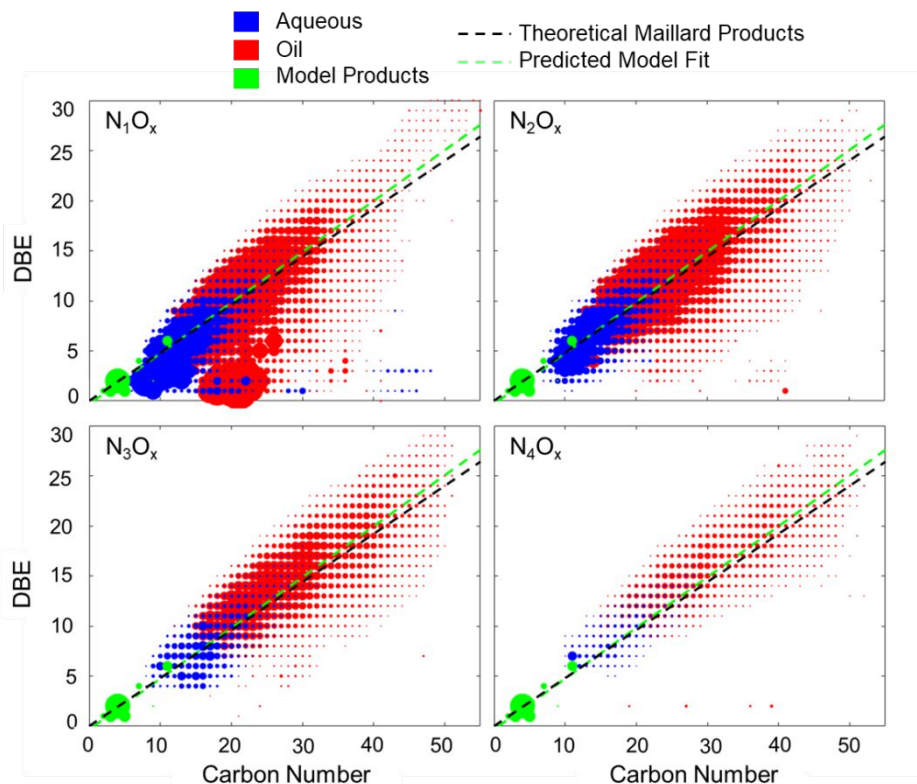


Figure 7. Double bond equivalency (DBE) versus carbon number for four nitrogen-containing heteroatom classes in the biocrude oil (red) and aqueous (blue) phases on food waste HTL at 575 K for 30 minutes. The green dots represented the products from the kinetic model (18% glycine, 22% diene, and 59% aldehyde split evenly between the four starting aldehydes). Model feedstock concentration was chosen to match that of the food waste HTL experiments. Experimental data points are sized by relative abundance and model points by relative concentration. The black dashed line represents the average trend for Maillard reaction products where the green line is the weighted average fit observed for products in the kinetic model.

To assess model robustness, a sensitivity analysis was performed to measure the effect of a 1% perturbation in rate constants on individual product yields. **Figure S4** reports these results, indicating the overall sensitivity of the competitive reactions 2.1, 3.1, 4.1, and 5.1 on overall product distribution and yields. The analysis highlights the competitiveness of the initial amine-oxygenate reactions and suggests that no single aldehyde functionality is solely responsible for the coupling reactions. Further discussion can be found in the Supporting Information.

Water also plays an integral role in the hydrothermal processing and chemical transformations examined in this study. In addition to being a reactant or product in 10 of the 38 reaction pathways examined, its ability to solvate the reaction species can play a potentially significant role in the reaction energetics and kinetics. The sensitivity analysis (**Figure S4**) confirms that water is in excess in the system and its final amount does not depend on the reaction rates ($\sigma_{ij} = 0$ for all instances of $i=\text{water}$). Future work should attempt to resolve the specific role

of water as a solvent in these key reaction pathways. In this study, all activity coefficients are assumed to be unity, indicating that non-ideal solvent interactions have been neglected. It has been shown that under supercritical conditions (675 K, 275 bar) the activity coefficient of water decreases to 0.5, thereby reducing the corresponding reaction rates by a factor of two.⁸³ While typical HTL reaction conditions are nearly 375 K and 75 bar below the explored supercritical conditions, the activity of water at these conditions remains relevant in improving the accuracy of the model.

The model pathways presented in this work are representative of elementary reactions that occur in the condensed phase chemistry during food waste HTL. As revealed by FT-ICR-MS, however, tens of thousands of unique species are detected across a wide molecular weight distributions ranging from light oxygenates to non-volatile hydrochars. Future work should 1) continue to explore elementary pathways with a larger breadth of functionality and sizes, 2) introduce solvent interactions to improve computational energetic calculation accuracy in the hydrothermal environment, and 3) consider the transient environmental changes in reaction environment due to primary (biopolymer degradation) and tertiary (polymerization) reactions.

Conclusions

Experimental analysis of food waste HTL products has revealed the presence of organic nitrogen in the form of fatty amides, indoles, and amines. Particularly notable is the prevalence of nitrogen-containing heterocyclic aromatics, the removal of which by conventional hydrodenitrogenation is difficult but necessary to hit fuel quality standards and avoid NO_x emissions. A detailed kinetic mechanism was proposed using model compounds representative of typical HTL feedstock monomers; subsequent DFT calculations and kinetic models were used to evaluate likely reaction pathways. The analysis revealed the Maillard reaction as a likely source of nitrogen heterocycle formation at HTL reaction conditions. FT-ICR MS characterization and analysis further supported this conclusion, revealing strong correlations between theoretical Maillard reaction products, experimentally measured HTL reaction products, and the model-predicted products.

These analyses serve to illuminate dominant chemistries and kinetic pathways by examining the activity of the highly reactive short chain intermediates that typically react on the order of seconds in HTL reactors. Substantial energetic barriers were shown to exist in deamination

and cyclization reactions, whereas condensation and ADA pathways experienced relatively low activation barriers, due to the relative stability of the transition states. This led to nitrogen heterocycles and gaseous nitrogen only being observed at higher temperatures; hemiaminals are the dominant product at lower temperatures. Strong compositional effects were observed in the formation of nitrogen heterocycles, which required stoichiometric amounts of amine, aldehyde, and diene. Sub-stoichiometric amounts of diene promoted production of hemiaminal and amide products, which is consistent with known hydrothermal products resulting from low lipid feedstocks. The predicted model trends correlated well with the experimentally observed product distributions from food waste HTL, with the FT-ICR MS showing final liquid products — dimer- and trimerization of amino acids, carbohydrate hydrolysis products, and fats — consistent with the Maillard reactions observed in the kinetic model. Ultimately, imines and aldehydes have been shown to be key classes in the proposed model, further showcasing its importance and accuracy in predicting HTL nitrogen intermediates. Future model expansions should consider the secondary reactions between the intermediates toward the formation of larger final products and the inclusion of solvent effects. Based on this work, the thermal barriers to deamination are revealed to be substantial kinetic limitations, so catalytic promotion of those pathways may be necessary to overcome large activation energetics.

Supporting Information

Supporting details can be found in the online version, including GC-MS, FT-ICR MS, and GC×GC methodology. Sensitivity analysis calculations and results, as well as molecule names and chemical classes for all species used in the model are also located in the Supporting Information. All FT-ICR MS spectra are provided via Open Science Framework <https://osf.io/64bmt/> through DOI 10.17605/OSF.IO/64BMT.

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