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Catalytic conversion of chitin as a nitrogen-containing biomass

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The conversion of chitin enables the utilisation of naturally-fixed nitrogen in addition to carbon toward establishing a sustainable carbon and nitrogen cycle. Chitin is an abundant biomass, 100 Gt per year, but most chitin-containing waste is discarded due to its recalcitrant properties. This feature article summarises the challenges and our work on chitin conversion to *N*-acetylglucosamine and oligomers with fascinating applications. Afterwards, we introduce recent progress on the chemical transformation of *N*-acetylglucosamine, followed by a discussion of future perspectives based on current status and findings.

Introduction

The basic concept of biomass utilisation is to establish a sustainable energy and material cycle. Current societies rely on fossil resources, but these fuels cannot regenerate within a reasonable timespan. Human activity required 6.4×10^{17} kJ of energy and released 36 Gt of CO₂ in 2021, *ca.* 20-fold larger than the amounts in 1900. The increased amount of CO₂ is causing the greenhouse effect and ocean acidification.¹ Meanwhile,

plants fix solar energy as organic materials using atmospheric CO₂, and the net primary production (NPP) is about 10^{17} carbon-g per year, accumulating 4×10^{18} kJ of energy.² Accordingly, biomass energy is several times larger than our demand. The utilisation of biomass, together with other renewable energy and material cycle systems, including artificial photosynthesis and chemical recycling of plastics,³ will become an indispensable technology for building a sustainable society.

Biomass conversion has mainly dealt with lignocellulose, starch and lipids, which are all composed of C, H and O atoms.^{4,5} Recently, artificially introducing N atoms to biomass-derived molecules has attracted significant interest as organic nitrogen compounds show a wide variety of functions.^{6–8} In this sense, chitin and chitosan, which are the most abundant biomass (annually *ca.* 100 Gt in total) next to lignocellulose,⁹ originally contain N atoms. Chitin and chitosan are both copolymers

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Fig. 1 Structures of chitin and chitosan. NAG and glucosamine units are randomly mixed. Chitin: $n/(n+m) > 40\%$, chitosan: $n/(n+m) \leq 40\%$ as a typical standard.

consisting of *N*-acetylglucosamine (NAG) and glucosamine units linked by β -1,4-glycosidic bonds (Fig. 1). Chitin contains more than 40 mol% NAG, and the polymers having lower ratios of NAG are called chitosan. Therefore, the conversion of chitin and chitosan enables the utilisation of naturally-fixed nitrogen to produce functional chemicals (Fig. 2).^{10,11} In terms of 'planetary boundaries', the utilisation of artificially-fixed nitrogen has exceeded the capacity of the Earth.¹² Thus, the nitrogen cycle is important in addition to carbon, and a potential solution is to use naturally-fixed nitrogen with high efficiency. Proteins also consist of nitrogen-containing monomers, but the structures are often complicated. The simple structures of chitin and chitosan may facilitate their utilisation in chemical industries.

While chitin and chitosan are attractive resources, they are resistant to degradation reactions, as described in detail in the following section. The development of new catalytic systems is crucial to achieving efficient utilisation of nitrogen-containing biomass.^{13–15} The selective scission of rigid chemical bonds within biomass molecules is especially important. Herein, this article summarises our recent work on the catalytic conversion of chitin to value-added organonitrogen compounds. We have preferred chitin over chitosan because the protection of reactive amino groups by acetyl groups is beneficial for selective chemical transformation. We show that acid catalysts including mineral acids and activated carbons hydrolyse chitin in the presence of mechanical milling. The section also discusses the reaction mechanisms and a significant application of chitin-



Fig. 2 Nitrogen-containing biorefinery.

oligosaccharides, the intermediates of hydrolysis. Afterwards, the authors introduce the conversion of NAG to useful derivatives that contain N atoms and its challenges, followed by concluding remarks.

Hydrolysis of chitin

Manufacture of NAG

Chitin contains many inter- and intra-molecular hydrogen bonds, which makes the polymer recalcitrant and insoluble in water. This nature is similar to that of cellulose, but there is a major difference between the two compounds. Cellulose forms hydrogen bonds to make two-dimensional sheets and they stack by dispersion forces, while chitin contains additional strong hydrogen bonds using amide groups (Fig. 3) to produce a three-dimensional hydrogen bond network.¹⁶ Even after the ball-milling treatment, chitin preserves most of the hydrogen bonds.¹⁷ The hydrolysis of cellulose is a difficult subject,¹⁸ but that of chitin requires even harsher reaction conditions. Moreover, chitin has amide bonds in addition to glycosidic bonds, and both can be hydrolysed. The selective scission of the glycosidic linkage is needed to synthesise NAG. Accordingly, the hydrolysis of chitin is a challenge.

To manufacture NAG from chitin, typical industrial processes use concentrated HCl and enzymes.^{19,20} A high concentration of



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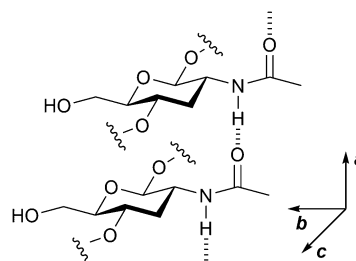


Fig. 3 Hydrogen bonds made by amide groups in chitin. Amide groups form hydrogen bonds along the *a*-axis direction. The image omits some functional groups for better visibility.



HCl is needed to accelerate the hydrolysis of glycosidic bonds while limiting deacetylation.^{21–23} The turnover numbers of acids are far less than 1 and this process produces a large amount of neutralisation waste. The hydrolysis of glycosidic bonds is an S_N1 reaction, and the bond cleavage after the pre-equilibrium of protonation is the rate-determining step (Fig. 4). Therefore, the hydrolysis rate relies strongly on the activity of H_3O^+ . In contrast, the hydrolysis of amide bonds under acidic conditions is an S_N2 reaction. The carbonyl oxygen in the amide structure is basic, easily undergoing protonation. Our density functional theory (DFT) calculations estimate that the proton affinities ($-\Delta H^\circ$ at 298 K) of the O atoms in amide and glycosidic bonds are 885 and 811 kJ mol⁻¹, respectively. Moreover, the rate-determining step is the addition of water.²⁴ Thus, the acid strength only slightly affects the reaction rate and the decrease in water activity at a high acid concentration slows down the reaction.^{21,24} Perhaps for a similar reason, pure formic acid selectively converts chitin to NAG formates with the retention of acetyl groups.²⁵ In contrast, a mixture of aprotic polar solvents and water accelerates deacetylation,²⁶ which is likely to be due to the increased nucleophilicity of water molecules by hindering hydrogen bonds ($H_2O \cdots H_2O$). The activity and nucleophilicity of water are major factors in determining the hydrolysis selectivity. Another method to synthesise NAG is acetylating glucosamine using acetic anhydride.²⁷ Water and methanol solvents allow the acetylation of amino groups while inhibiting the esterification of hydroxy groups. However, the direct synthesis of NAG from chitin would be better. These factors have pushed the price of NAG to over 50 GBP per kilogram. An efficient process for hydrolysing chitin to NAG is essential to achieve a chitin-based biorefinery.

Hydrolysis of chitin by mineral acids under ball-milling conditions

Dilute mineral acids are ineffective for the selective hydrolysis of chitin as described above. Among other types of catalyst, carbon materials with weak acid sites characteristically hydrolyse cellulose in water,^{18,28} but this system does not function for chitin as it is. Chitin is more recalcitrant than cellulose, requiring us to add a new idea for achieving its hydrolysis.

Previously, several groups have reported that acid catalysts hydrolyse cellulose to oligomers during the milling process.^{29–33}

We expected that the mechanical forces of ball-milling will activate the recalcitrant chitin, thereby allowing the hydrolysis of chitin with a small amount of acid.³⁴ Indeed, the effect of milling was obvious; a catalytic amount of H_2SO_4 quantitatively hydrolysed chitin to soluble compounds under dry planetary milling (Fig. 5). In detail, we added H_2SO_4 to chitin powder (substrate/catalyst ratio (S/C) = 8.1 mol-NAG unit/mol- H_2SO_4) dispersed in diethylether, and the sample was dried by a rotary evaporator. The resulting powder was milled using alumina balls and an alumina pot at 500 rpm for 6 h. The temperature inside the pot was at most 60 °C during milling. No solvent was used for this reaction, but physisorbed water on the sample (2–5 wt%) worked as a reagent for hydrolysis. ¹H nuclear magnetic resonance (NMR) spectroscopy showed that the major products were oligomers of NAG with β -1,4-glycosidic bonds and 1,6-bonds including anhydride structures. Small quantities of unidentified products were also included. Surprisingly, acetic acid (AcOH) was not detected in the product mixture, suggesting no hydrolysis of amide bonds for the following reasons: first, low water content is beneficial to reducing deacetylation (Fig. 4). Second, the mechanical forces selectively accelerate the hydrolysis of glycosidic bonds (see below). Therefore, the reaction cleaved glycosidic bonds while preserving amide bonds.

The solid sample produced by milling with H_2SO_4 underwent further hydrolysis in water solvent at 190 °C, giving NAG in 32% yield, based on original chitin.³⁴ The yield increased to 61% at an S/C of 3.4 at 175 °C (Fig. 6).³⁵ The turnover number of H_2SO_4 for the formation of NAG was 2.1, which was a catalytic reaction.

Instead of hydrolysis, alcoholysis is also possible for degrading the oligomer intermediates (Fig. 6). Changing the solvent to methanol, we synthesised 1-O-methylated NAG in up to 70% yield.³⁴ NAG is an unstable compound, but alcoholysis can protect the reactive hemiacetal groups, decreasing the decomposition of products.³⁶

Mechanistic study of mechanocatalytic hydrolysis

To clarify the effect of milling, we designed several hydrolysis experiments and measured the solubilisation ratios of the products by preparing a 1 wt% suspension in water at room temperature.³⁴ Initially, we confirmed that chitin contains no



Fig. 4 Hydrolysis mechanisms of glycosidic and amide bonds. The image omits some functional groups for better visibility.



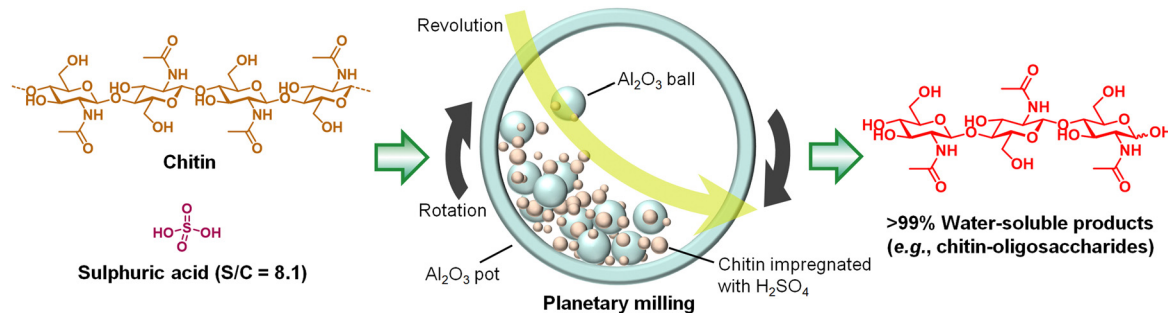


Fig. 5 Hydrolysis of chitin by H₂SO₄ in the mechanochemical method.

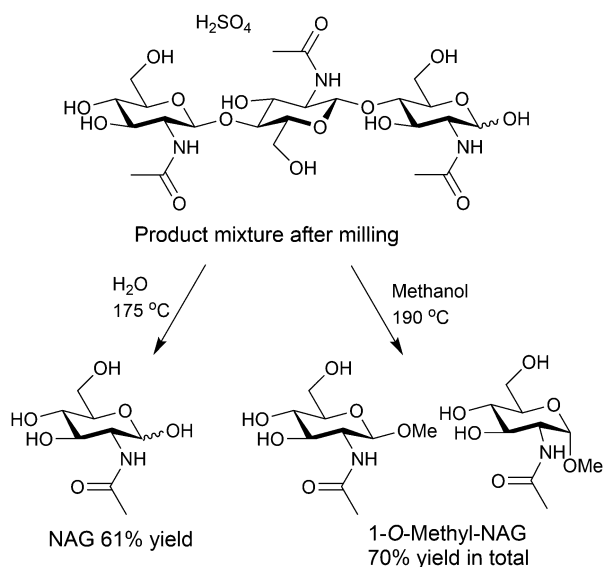


Fig. 6 Conversion of the oligomers produced by a mechanochemical reaction to monomers.

soluble products. In the second experiment, chitin was milled for 6 h, and the product showed a solubilisation ratio of 5.2%. Accordingly, the milling treatment causes a very limited degree of hydrolysis in the absence of a catalyst. For the next control experiment, chitin was milled, impregnated with H₂SO₄ and aged for 6 h, which showed a solubilisation ratio of only 7.8%. H₂SO₄ alone does not hydrolyse chitin. By altering the order, chitin was impregnated with H₂SO₄ and milled for 6 h, giving >99% solubilisation ratio. These results show that H₂SO₄ hydrolyses chitin under milling conditions.

Acid strength dominates the solubilisation ratio after the milling process.³⁵ Chitin was impregnated with several acids with different pK_a values, and the samples were milled for 6 h. HClO₄ (pK_a = -10) and H₂SO₄ (pK_a = -3) gave completely soluble products. CH₃SO₃H (pK_a = -2) and C₃F₇CO₂H (pK_a = 0.4) showed ca. 90% solubilisation, followed by H₃PO₄ (pK_a = 2.1) (44% solubilisation) and AcOH (pK_a = 4.7) (2.8% solubilisation). These results suggest that the protonation of glycosidic bonds is an essential step for hydrolysis under our milling conditions.

In the planetary milling process, alumina balls accelerated by the centrifugal force hit the pot wall. This event provides

mechanical forces and heat at the collision point. We performed control experiments to reveal which factor is predominant in facilitating the hydrolysis of chitin (Fig. 7).³⁷ A planetary milling of H₃PO₄-impregnated chitin afforded a pale yellow product containing NAG and oligomers consisting of two to six NAG units in 40% yield in total. This reaction produced no AcOH. In control experiments, chitin was milled, impregnated with H₃PO₄ and aged with no solvent at a high temperature in the range of 60–120 °C for a designated time. The reaction temperature was decided by the following assumption; the temperature of the bulk phase was at most 60 °C during the milling and the increase in the temperature at the local area by ball collision was estimated to be roughly as high as 50 °C. The optimum reaction produced NAG and oligomers in 33% yield. However, the product was dark brown and contained AcOH in 24 mol% yield based on the number of NAG units in chitin. Accordingly, the effect of milling cannot be ascribed to local heat but to mechanical forces. This is a mechanocatalytic reaction.

We speculated that mechanical forces easily work along the polymer chain of a chitin molecule, while they are rarely available within side-chains, namely amide groups.³⁴ However, it was unclear how great forces are applied to chitin during milling and if they are large enough to activate glycosidic bonds

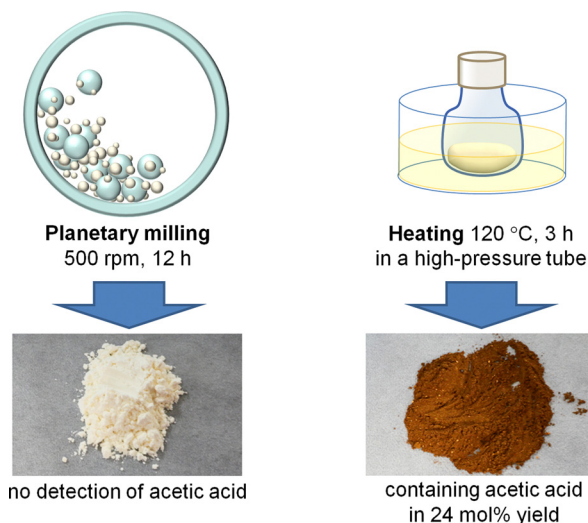


Fig. 7 Comparison of milling and heating reactions. Chitin was impregnated with H₃PO₄ (S/C = 5.0) before the treatments.



in chitin. These questions are also applicable to cellulose.^{29–33} In a simulation of milling, Rinaldi *et al.* and Takagaki *et al.*, respectively, estimated the relationship between the total milling energy and hydrolysis rate, which showed good correlation.^{32,38} To consider the effect of mechanical forces, we solved the motion equation of an alumina ball during planetary milling with a rough approximation, which indicated that the alumina ball hits the pot wall at a velocity of 4 m s^{-1} .³⁷ Based on the collision velocity, finite element simulations estimated that up to 7 GPa of compressive stress and 1.5 GPa of tensile stress are imposed on chitin and cellulose. Therefore, the compressive stress is larger, except when stress concentration occurs. Multiplying by the molecular cross-sectional area, we can estimate that the maximum tensile and compressive stresses correspond to 0.5 and 2.4 nN of forces, respectively, on one chitin or cellulose molecule. These values are similar to the connecting forces of covalent bonds, as evaluated by pulling polymer molecules with atomic force microscopes.³⁹

DFT calculations were employed to estimate the effects of forces on chemical bonds. Amirjalayer *et al.* showed that tensile forces decrease the activation energy of hydrolysing glycosidic bonds.⁴⁰ Using the B3LYP functional^{41,42} within DFT, we demonstrated that the conformation of the tetrahydropyran ring of NAG changes to increase the electron density on C–O σ^* bonds in the glycosidic bonds under a tensile-force field.⁴³ This facilitates an elimination reaction to cleave the glycosidic bonds in a *syn* form between an O lone pair and C–O bond (Fig. 8). This type of reaction is rare as the *syn* orientation provides larger steric repulsion due to the eclipsed conformation and a smaller overlap of orbitals than the *anti* forms. This indicates that forces could alter reaction mechanisms from the usual thermal reactions. In addition, our calculations showed that the tensile forces do not accelerate the hydrolysis of amide.⁴³ The rate-determining step of acid-catalysed amide

hydrolysis is the addition of water (Fig. 4), which is not related to the tensile forces on the amide group. This feature is different from alkaline hydrolysis under a mechanochemical condition that cleaves amide bonds,⁴⁴ where the scission of C–N bonds is the rate-determining step for amide hydrolysis.⁴⁵

We have also demonstrated that not only tensile forces but also compressive ones activate glycosidic bonds in hydrolysis.³⁷ Cellobiose was employed as a small model to reproduce the effects of forces, which significantly reduced the calculation time. In the DFT calculations at the B3LYP/6-31+G(d,p) level, the conformation of H_3PO_4 -coordinated cellobiose was initially optimised, and then the distance between two oxygen atoms at the tips of the molecule was changed to provide tensile or compressive forces. Then, we simulated hydrolysis reactions in the force-affected molecules (Fig. 9). This procedure, called the COGEF method,^{46,47} slightly changes the intensity of forces during reactions but easily evaluates the effect of forces. Note that the protonation of cellobiose is simpler than the coordination of H_3PO_4 , but cellobiose protonated at the glycosidic bond underwent bond scission with only 7.4 kJ mol^{-1} of activation energy. The naked proton is too active and it does not reproduce actual experiments. In the absence of forces, the glycosidic bond of H_3PO_4 -coordinated cellobiose was cleaved at an activation energy of 120 kJ mol^{-1} . An anti-periplanar conformation appeared in the transition state, which is similar to typical E2 reactions. Stretching the cellobiose molecule by 7.5%, corresponding to *ca.* 3 nN of pulling force, decreased the energy barrier to 93 kJ mol^{-1} . The energy needed can be further decreased to 76 kJ mol^{-1} at 9.4% elongation of the molecule.

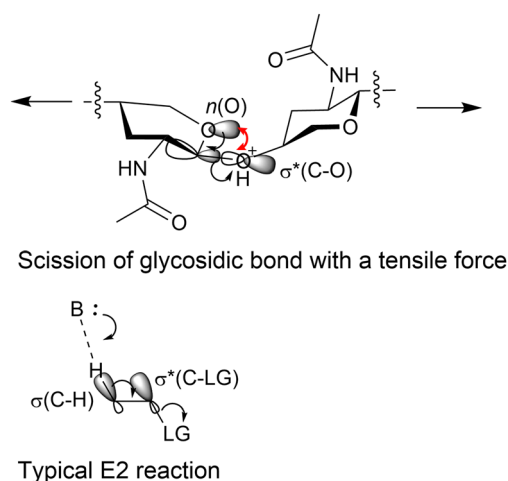


Fig. 8 Reaction mechanisms of bond cleavage. The image omits some functional groups of chitin for better visibility. In the *syn* orientation (top), the red arrow highlights opposite signs between the $n(\text{O})$ and the OH side of $\sigma^*(\text{C}-\text{O})$ orbitals. In contrast, the *anti* orientation (bottom) does not have such a repulsion.

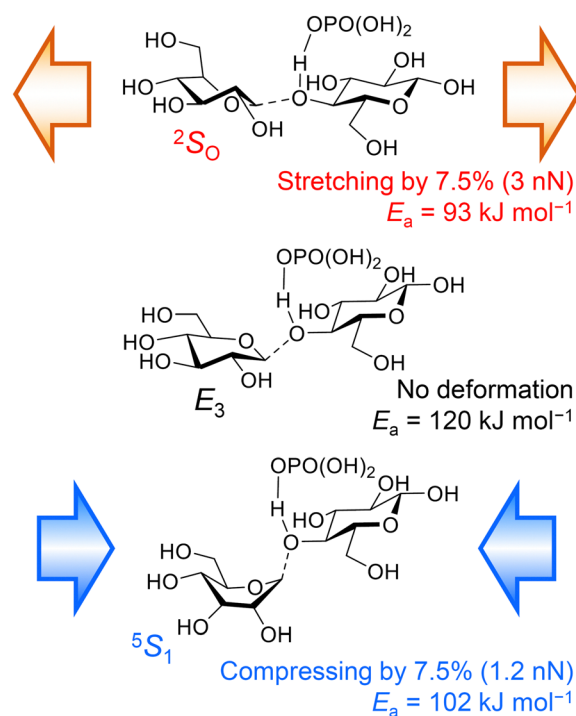


Fig. 9 Effects of forces on the cleavage of glycosidic bonds. The E_a value is not an activation energy under a constant force but under a constant distance between the tips of the molecule.



A *syn* conformation was found during the bond cleavage, as described above. Meanwhile, compressing by 7.5%, *ca.* 1.2 nN of force, also lowered the required energy to 102 kJ mol⁻¹. The force stabilised the twisted boat structure, giving an anti-periplanar conformation for the bond scission. It is easy to assume that a tensile force activates a chemical bond, but our calculations show that a compressive force also works to some extent. Moreover, our classical mechanics calculations have indicated that the maximum tensile and compressive forces are 0.5 and 2.4 nN, as described above. Taking this fact into account, the compressive forces produced by planetary milling are large enough to activate glycosidic bonds, whereas tensile forces can do so only when there is force concentration. Thus, we propose that compressive forces are as important as tensile forces in the mechanocatalytic reaction.

DFT is designed to calculate the ground states of molecular systems.⁴⁸ However, in the presence of tensile forces, chemical bonds are stretched in the homolysis manner, which may mix excited states. To evaluate the accuracy of the DFT calculations, we compared B3LYP and the full configuration interaction (full CI) with the aug-cc-pV5Z basis set^{49,50} and the closed shell system in terms of the energy needed to stretch H₂.³⁷ Surprisingly, their difference was less than 0.5 kJ mol⁻¹ up to 12% elongation. Stretching the CO molecule also gave similar energies between B3LYP and the complete active-space second-order perturbation theory (CASPT2; ten electrons, eight orbitals)⁵¹ up to 25% elongation. Hence, the calculation method is reliable over this range.

Catalysis of oxygenated carbon in the hydrolysis

Mineral acids are active for the mechanocatalytic hydrolysis of chitin, but enabling the use of solid catalysts may expand practical applications. Kerton *et al.* reported that kaolinite, an acidic layered clay mineral, can accelerate the hydrolysis of chitin in the milling process.⁵² Later, we found that carbons with weak acid sites are active and selective catalysts for converting chitin to chitin-oligosaccharides under planetary milling conditions.⁵³ This is different from the conventional aqueous-phase reaction, where they do not hydrolyse milled chitin, as described above.

In our work, a carbon-based catalyst was prepared by the air oxidation^{54,55} of a commercial activated carbon (BA) at 425 °C. The recovery of the catalyst, which was named AC-Air, was about 50 wt%, and the remaining part was burnt out. AC-Air had 0.90 mmol g⁻¹ of carboxylic groups and 0.84 mmol g⁻¹ of phenolic hydroxy groups. Chitin was treated with AC-Air in a planetary mill (Fritsch P-6) at 500 rpm for 12 h. The product showed a 34% solubilisation ratio and contained water-soluble oligosaccharides in 32% yield (>90% selectivity) (Fig. 10). In contrast, the reaction in the absence of a catalyst gave an 8% solubilisation ratio and produced oligosaccharides in 1.2% yield. Accordingly, the catalyst elevated the yield of oligosaccharides by 27 times, compared to the non-catalytic reaction. This enhancement is remarkable as kaolinite increased the yield by only three times in previous work.⁵² In the presence of AC-Air, extending the milling time to 48 h increased the solubilisation ratio to 72% and the yield of oligosaccharides to 66%. Accordingly, this reaction

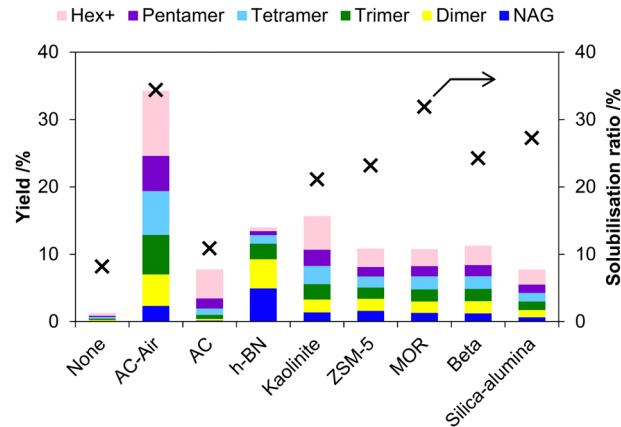


Fig. 10 Screening of solid catalysts in the mechanocatalytic hydrolysis of chitin. Hex+ indicates the total yield of hexamer and larger oligomers. Yield is based on moles of carbon.

system provides more than 90% selectivity for oligosaccharides among soluble products (66/72 = 92%).

¹H NMR analysis showed a high purity of chitin-oligosaccharides in the water-soluble product after 48 h of milling with AC-Air. We focused on the peaks of C1-H (Fig. 11) as they specifically appear in a low magnetic field region. The major peaks were assignable to chitin-oligosaccharides, namely β-1,4-oligomers of NAG. Additionally, anhydrides were found, but the peak area was only about 5% among all C1-H peaks. In contrast, when we used H₂SO₄ as a catalyst, the product gave many peaks in the C1-H region, suggesting the presence of oligomers with different structures.

The selective production of chitin-oligosaccharides has a fascinating application in agriculture. Recently, it has been a challenge to decrease the use of agrichemicals to retain high productivity while preserving the natural environment. To overcome this problem, biostimulants that bring out the abilities of plants themselves will be an alternative to conventional agrichemicals,⁵⁶ and the market will grow up to 6–8 billion GBP per year by 2030. In this sense, chitin-oligosaccharides can activate plant immune systems.^{57,58} If plants are infected with fungi, oligosaccharides present on the microbe cells are released (Fig. 12). Plants detect the fragments as pathogen-associated molecular patterns (PAMPs) and express immune-related genes. Relatively large but water-soluble chitin-oligosaccharides work as PAMPs so that plants become more resistant to microbes. Additionally, oligosaccharide mixtures containing chitin-oligosaccharides assist the growth of plants, possibly increasing the yield of crops.⁵⁹

Taking into account the application of chitin-oligosaccharides, the distribution of polymerisation degree is of interest.

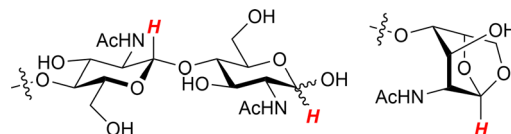


Fig. 11 C1-H (highlighted in red) in oligomers and anhydrides.



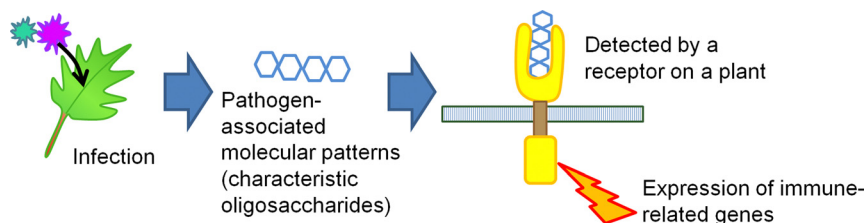


Fig. 12 Schematic of PAMP recognition.

We changed the reaction time from 12 h to 48 h using AC-Air. Major products were trimer to hexamer without dependence on the reaction time, and the successive hydrolysis was limited. This result is noteworthy, because small oligomers are usually more reactive than large molecules, giving monomers as the predominant product and only a small amount of oligomers.^{60,61} As a representative example, a trial using 11 M HCl to convert chitin to oligosaccharides produced NAG as a major product and the yield of oligomers decreased with increasing polymerisation degree.⁶² Our Monte-Carlo simulations indicated that AC-Air cleaves glycosidic bonds almost with the same probability regardless of the polymerisation degree. Therefore, for example, a decamer (with nine glycosidic bonds) undergoes hydrolysis roughly nine times faster than a dimer (one glycosidic bond). These kinetics allow for the accumulation of a water-soluble but not too small range of chitin-oligosaccharides, namely bioactive ones. Carbons can easily adsorb large polysaccharides,⁶³ and the homogeneously adsorbed polymeric substrates have the chance to receive mechanical forces. This system is suited for synthesising chitin-oligosaccharides working as PAMPs.

We further elucidated the cause of the high activity of AC-Air by examining related acid catalysts (Fig. 10). While AC-Air gave 32% yield of oligosaccharides after 12 h of milling, pristine activated carbon (AC) was inactive (oligosaccharides 6.2% yield), suggesting that the oxygenated functional groups with weak acidity are the active sites. Hexagonal boron nitride, with a similar structure to graphite and very low acid strength,^{64,65} gave 9.1% yield of oligosaccharides. Kaolinite was the second most active (14% yield) among the catalysts we tested. Proton-type zeolites (ZSM-5, beta and MOR) and silica-alumina produced oligosaccharides in 7–10% yields. A sulfonic acid resin, Amberlyst, almost completely converted chitin to soluble products. However, Amberlyst became soluble in water after the reaction, suggesting

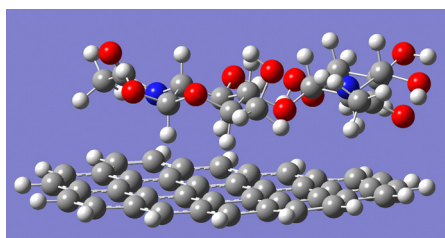


Fig. 13 Adsorption structure of *N,N'*-diacetylchitobiose on carbon. Optimised at B3LYP-D3/6-31G(d,p). Many C–H groups of the sugar are directed toward aromatic rings, and the carbon is slightly bent to fit the sugar molecule.

that this resin was virtually similar to H_2SO_4 in this system. Accordingly, a solid catalyst should have many acid sites but could be a weak acid such as carboxylic acid. Moreover, layered materials, especially carbons, are better than typical three-dimensionally bonded oxides. In the mechanocatalytic hydrolysis of cellulose, Takagaki *et al.* have shown the activity of HNbMoO_6 ,³² which is also a layered material. We guess that the thin-layer structure is useful to contact with chitin particles at the molecular level due to its flexibility. In addition, a carbon material has polycyclic aromatics as the basal plane, and they attract chitin by $\text{CH}-\pi$ interactions (Fig. 13).^{65–67} This is different from the fact that oxides apply electrostatic interactions. These characteristics make the air-oxidised carbon more active for the hydrolysis of chitin.

Conversion of NAG

Potential derivatives of NAG

NAG is a potential feedstock to produce a variety of chemicals (Fig. 14). Thermal decomposition of NAG or chitin produces many compounds, such as pyridine, pyrazine, chromogens, 3-acetamido-5-acetylfuran (3A5AF) and 3-acetamidofuran.^{68–71} The use of NaOH can change the product distribution, giving a small amount of pyrrole.⁷² An important and characteristic product among the derivatives is 3A5AF. This compound can be a precursor to proximicins, which are potential anticancer medicines.⁷³ The oxidation of 3A5AF may produce 3-acetamido-5-carboxyfuran (3A5CF), with one carboxy and one amide group, likely to be a monomer of polyamide. Reacting 3A5AF with a methyl Grignard reagent, followed by dehydration, leads to vinyl compounds.⁷⁴ Its radical polymerisation perhaps produces side-chain polymers. Meanwhile, the hydrogenation of NAG selectively produces 2-acetamido-2-deoxysorbitol (ADS).^{75,76} This compound is transformed into 2-acetamido-2-deoxyisorbide (ADI), a promising precursor to polymers.⁷⁷ An oxazolidinone with 5-(*S*) configuration, a potential precursor to antibacterial agents, is also synthesised from ADS.⁷⁸ Combining a retroaldol reaction and hydrogenation converts NAG to acetyl monoethanolamine, and its oxidation provides an amino acid in a protected form.⁷⁹ The oxidation of NAG also produces an amino acid, specifically *N*-acetylglucosaminic acid.⁸⁰ Supported Au catalysts convert NAG and related amino sugars such as mannosamine to corresponding amino acids. The system is applicable to chitosan for synthesising glucosaminic acid.⁸¹ As most studies on NAG conversion have been started in recent years, further efforts will expand the potential derivatives and applications. The following sections focus on 3A5AF and ADI as their applications are promising.



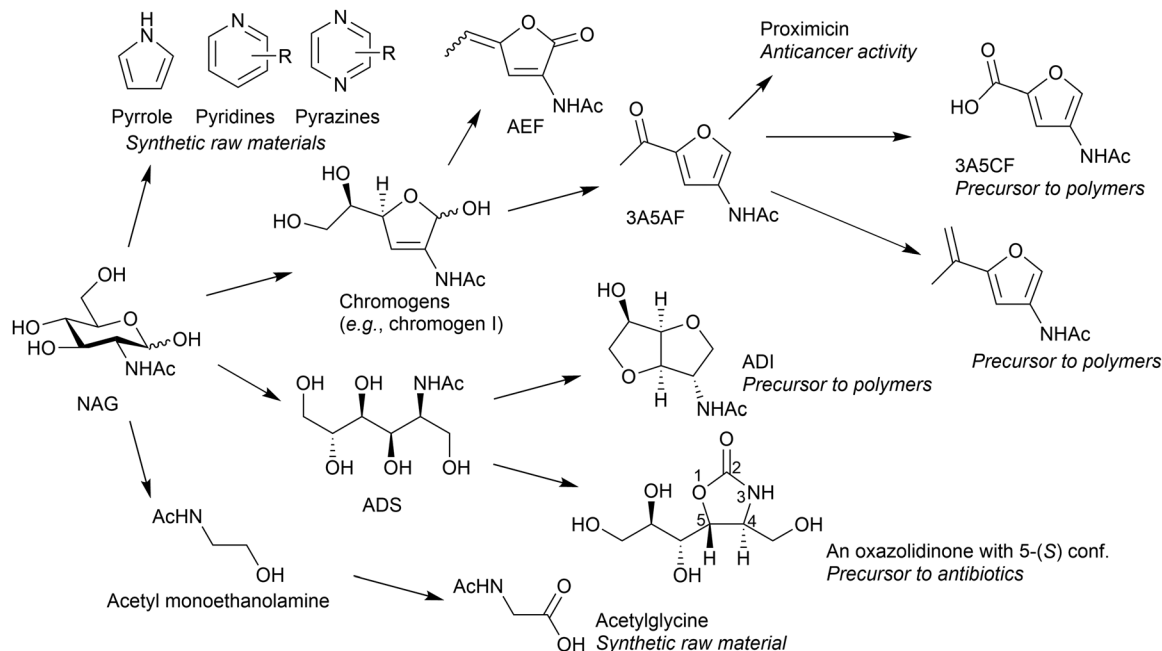


Fig. 14 Conversion of NAG to organonitrogen compounds and their potential applications.

Conversion of NAG to 3A5AF

The furanic compound 3A5AF was found in a small quantity in the pyrolysis products of NAG and chitin.^{68,69} Kerton *et al.* reported a liquid-phase conversion of NAG to 3A5AF in up to 60% yield in the presence of 1-butyl-3-methylimidazolium chloride ([BMIM]Cl) and B(OH)₃ under microwave irradiation (Table 1, entry 1).⁸² The ionic liquid can be replaced with dimethylacetamide (DMAc) and NaCl (entry 2).⁸³ Afterwards, Yan and co-workers found that B(OH)₃ and NaCl produce 3A5AF in 7.5% yield from chitin in *N*-methylpyrrolidone (NMP) (entry 3).⁸⁴ The use of ball-milled chitin improves the yield of 3A5AF to 29% (entry 4).⁸⁵ They also clarified that Cl⁻ significantly increases 3A5AF yield, by comparing several anions.⁸⁶

For the reaction mechanism, it is well known that a similar dehydration reaction of glucose produces 5-hydroxymethyl-2-furaldehyde (5-HMF). However, the positions of the functional groups in 3A5AF and 5-HMF are different (Fig. 15). Glucose initially isomerises to fructose, which is a ketose with a

furanose form, followed by dehydration (Fig. 15b). However, NAG cannot isomerise to the form corresponding to fructose due to a large disadvantage in energy (+73 kJ mol⁻¹, predicted with DFT calculations).⁷⁹ Therefore, NAG directly undergoes dehydration possibly in the linear form, giving functional groups at 3- and 5-positions (Fig. 15(a)), although the detailed reaction mechanism is under discussion.^{82,86–88} Boron compounds may shift the equilibrium among the pyranose, furanose and linear forms by producing esters with NAG toward accelerating dehydration. Cl⁻ might change the hydrogen bonding structures of sugars. Note that the deacetylation of NAG enables the formation of 5-HMF, because a double bond (C=N) can be produced in the resulting structure.^{89–92} Accordingly, the amide group plays a predominant role in the NAG transformation.

The above-mentioned systems use B(OH)₃, but its removal is sometimes problematic as B(OH)₃ easily produces adducts with sugar compounds. In addition, these reactions need high temperatures, often with unconventional methods. Decreasing the reaction temperature to 140 °C dropped the yield of 3A5AF to 1% despite the almost complete conversion of NAG in the presence of NaCl and B(OH)₃ in DMAc.⁹³ One probable reason is the low solubility of NaCl in the aprotic solvents.⁹⁴ Rapid heating to a high temperature dissolves the salt to accelerate the desired reaction before side-reactions.

Accordingly, we need an easy method that many researchers can use to synthesise and isolate 3A5AF for further studies.

Previous studies suggest that the conversion of NAG to 3A5AF needs aprotic amide solvents and chloride ions. B(OH)₃ could be a chelating agent or a Lewis acid. Based on these assumptions, we found that AlCl₃·6H₂O transforms NAG into 3A5AF in 30% yield in dimethylformamide (DMF) at 120 °C by conventional oil-bath heating without adding boron compounds (Table 1, entry 5).⁹³

Table 1 Synthetic systems of 3A5AF from NAG

Entry	Substrate	Catalyst and additive (mol%)	Solvent	T/°C	Yield of 3A5AF/%	Ref.
1	NAG	B(OH) ₃ (200)	[BMIM]Cl	180 ^a	60	82
2	NAG	B(OH) ₃ (100), NaCl (400)	DMAc	220 ^a	58	83
3	Chitin	B(OH) ₃ (400), NaCl (200)	NMP	215	7.5	84
4	Ball-milled chitin	B(OH) ₃ (400), HCl (100)	[BMIM]Cl	180	29	85
5	NAG	AlCl ₃ ·6H ₂ O (100)	DMF	120	30	93

^a Microwave heating.



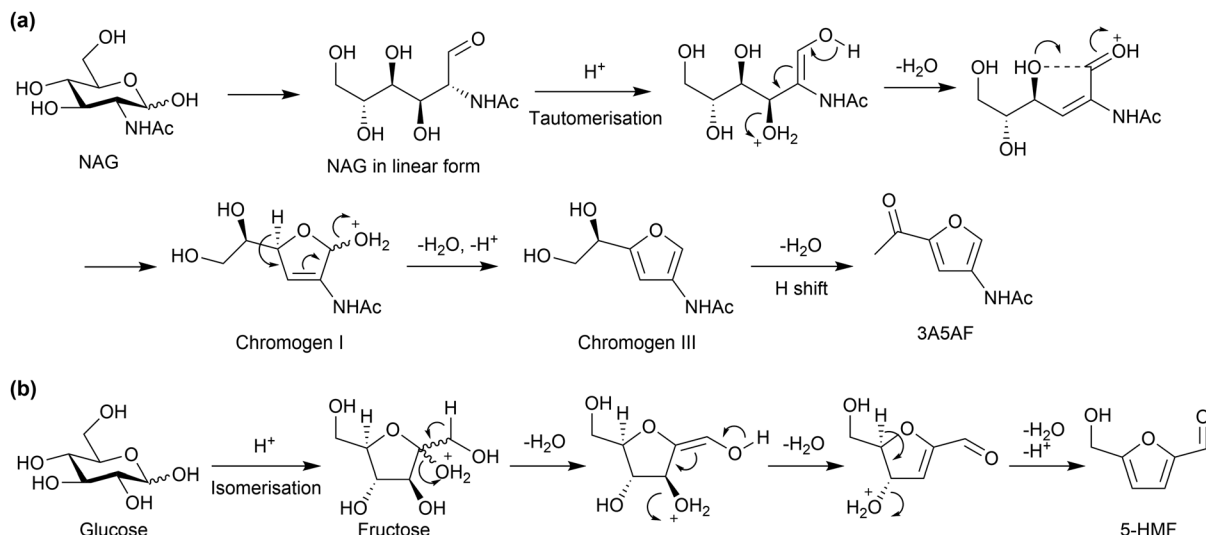


Fig. 15 Plausible reaction mechanisms to produce (a) 3A5AF and (b) 5-HMF. The dehydration reactions have derivatives in the mechanisms.

Moreover, this reaction can be performed at a 100 mL scale with 5% concentration of NAG, giving 0.70 g of 3A5AF (18% isolated yield) with more than 98% purity after silica gel chromatography. This reaction also generated 3-acetamido-5-ethylidene-2(5*H*)-furanone (AEF), with a sweet scent similar to dairy products, as a side product. The yield of AEF increased to 7% by using $\text{Al}(\text{OTf})_3$ as a catalyst. This compound could also be interesting.

In summary, 3A5AF is available at a laboratory scale, but the efficiency is not very high. We should further improve the synthetic method to make the compound more useful and easily accessible.

The oxidation of an acetyl group in 3A5AF produces 3A5CF, with one carboxy and one amide group. This compound is likely to be a promising precursor to polyamides. However, the conversion of 3A5AF to 3A5CF is a challenge. Currently, the only way to achieve this transformation is to use the iodoform reaction with a stoichiometric amount of iodine (Fig. 16(a)).⁷³ In other oxidation reactions, the Baeyer–Villiger oxidation of 3A5AF should produce a predominant isomer (Fig. 16(b)), while radical reactions such as autoxidation probably attack the furan ring due to the low aromatic stabilisation.^{95,96} It is desirable to

develop an efficient catalytic method to produce 3A5CF by overcoming these problems.

Synthesis of ADI

This condensed five-membered ring compound has one amide and one hydroxy group (Fig. 17(a)). Although the application has not yet been developed, the structure is attractive as a feedstock for polymer synthesis. Isosorbide, with the same skeleton as ADI and two hydroxy groups, is a precursor to engineering plastics (Fig. 17(b)).⁹⁷ Indeed, the isosorbide-based polycarbonate named Durabio shows excellent scratch resistance, high transparency with remarkable chromogenic property and high thermal resistance, and is thus applied in automotive parts (*e.g.*, Mazda CX-5) and mobile phones.⁹⁸ ADI-based plastics may also exhibit remarkable properties. ADI may be synthesised from NAG in a similar manner to the glucose-to-isosorbide system, and we reported the synthesis of ADI from NAG for the first time.⁷⁷ The production of an ADS intermediate has been established at a laboratory scale as follows: NAG (4 wt% concentration) was hydrogenated over Ru/C under 4 MPa of H_2 in water at 80 °C, and the product mixture was filtrated and evaporated. The resulting condensed material was dissolved in methanol at 60 °C and recrystallised at –30 °C, giving 85% isolated yield of ADS. The concentration of NAG can be increased to improve the productivity. The produced ADS was converted to ADI by a $\text{CF}_3\text{SO}_3\text{H}$ catalyst (*S/C* = 2) at 150 °C, which afforded 33% yield of ADI.

The conversion of ADS needs a large amount of $\text{CF}_3\text{SO}_3\text{H}$ (*S/C* = 2), unlike sorbitol undergoing dehydration with a small amount of H_2SO_4 or H-beta zeolite.^{99,100} A major reason is the high basicity of the amide group in ADS.⁷⁶ The carbonyl oxygen traps protons, which increases the energy for protonating hydroxy groups for the dehydration reaction. Our DFT calculations have shown that the second dehydration step of ADS is the rate-determining step, and the activation energy is 58 kJ mol^{-1} higher than the corresponding reaction of sorbitol. To decrease the influence of proton trapping, a large amount of strong acid



Fig. 16 (a) Iodoform reaction and (b) Baeyer–Villiger oxidation of 3A5AF.





Fig. 17 Similarity between (a) conversion of NAG to ADI and (b) that of glucose to isosorbide.

was needed. The addition of lanthanoid triflate slightly increases the yield of ADI to 40%, perhaps by coordinating with the amide group of the substrate to hamper proton trapping.¹⁰¹

We explored the possibility of replacing strong acids with weak acids and unexpectedly found that H₃PO₃ accelerates the conversion of ADS to ADI.¹⁰² Other acids with similar pK_a values (H₃PO₃, 1.3) showed much lower activity for this reaction. NMR analysis indicated the presence of phosphite esters of ADS and anhydro-ADS that are possible intermediates. To clarify the role of the esters, cyclisation reactions of the esters to produce ADI were simulated by DFT calculations. The second cyclisation reaction gave 82 kJ mol⁻¹ of activation energy for the phosphite ester system (Fig. 18 phosphite mechanism), which was 36 kJ mol⁻¹ lower than that in the absence of the ester (Fig. 18 acid mechanism). H₃PO₃ easily produces an ester with anhydro-ADS, and its P=O group receives a proton with a significantly lower energy than a hydroxy group for producing ADI. Although the yield of ADI has been at most 25%, this result indicates that the phosphite mechanism could overcome the proton trapping problem. Nozaki *et al.* have proposed that Al(PO₃)₃ produces esters with phenolic hydroxy groups, which facilitates the hydrodeoxygenation of the

phenolic compounds over a Pt/Al(PO₃)₃ catalyst.¹⁰³ The utilisation of esters is an interesting approach to activate hydroxy groups.

Conclusions and outlook

Chitin is a simple molecule consisting of nitrogen-containing sugars, and therefore it is a fascinating feedstock for producing organonitrogen compounds. However, the strong hydrogen bonds and the presence of multiple reactive groups in chitin have hampered its selective and efficient hydrolysis to NAG. We found that mechanical forces applied in planetary milling selectively activate glycosidic bonds so that acids can cleave the bonds while preserving amide bonds. Our quantitative analysis of mechanical forces indicated that tensile and compressive forces produced in the planetary mill both contribute to dissociating glycosidic bonds. Moreover, the forces could change the reaction mechanism. This mechanochemical system can utilise solid catalysts and a carbon-based weak acid is especially effective. The catalyst generates chitin-oligosaccharides in a high yield, and the product can not only be a source of NAG but also a

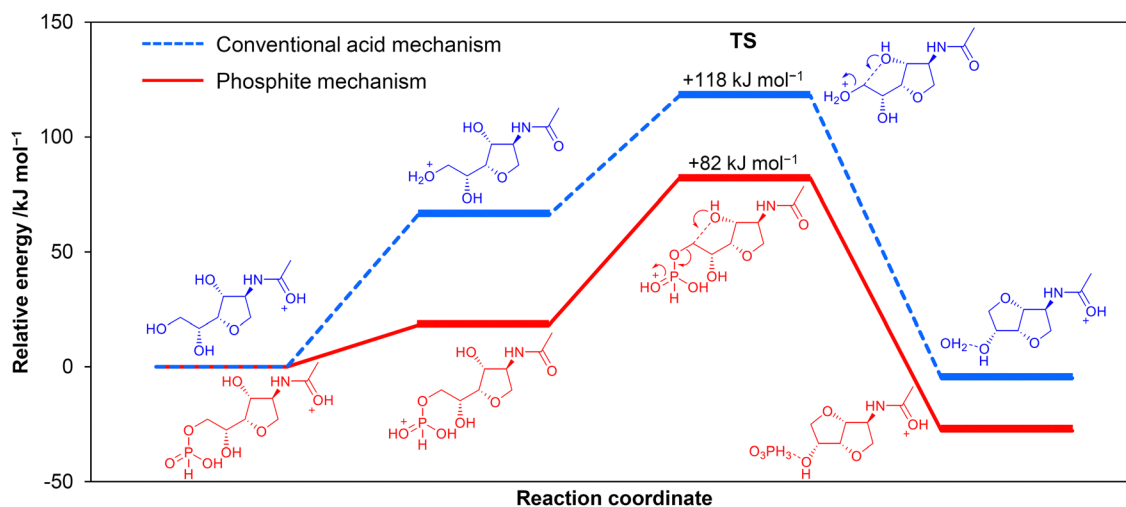


Fig. 18 Energy profiles for the cyclisation reactions to produce ADI, determined by DFT calculations. Red: phosphite mechanism. Blue: conventional acid mechanism. Calculation level: B3LYP/6-31+G(d,p) with solvent effects of DMF by SMD. The zero-point vibration energy was included. The energy required to transfer a proton from amide to P=O (phosphite mechanism) or hydroxy (acid mechanism) groups dominates the activation energy.



biostimulant that assists the growth of plants with a smaller amount of agrichemicals. The latter application provides a high value to chitin-oligosaccharides so it makes the implementation much easier than usual bulk chemical productions. The limitation of the current hydrolysis system is that it consumes a considerable amount of energy in the milling process during the long reaction time. Although the use of renewable energy may alleviate this issue, we hope that the development of new catalysts and mechanochemical systems based on our mechanistic studies will improve energy efficiency. In addition, the use of deep eutectic solvents or highly concentrated salt solutions has recently emerged as a potential method to activate chitin.^{104–107} They would also contribute to the selective extraction of chitin from biomass, because the current extraction method uses a large amount of NaOH and HCl.^{10,108} These technologies based on solid science will expand the possibility of chitin conversion. Moreover, as mentioned in a previous section, chitin is similar to but more recalcitrant than cellulose, so the chemistry established here may also be useful in cellulose conversion.

The synthesis of NAG derivatives is an emerging topic and has not been very efficient, except for ADS. The latest studies have clarified that we should pay attention to the different reactivity of the amide group from a hydroxy group (glucose). Amide has much higher basicity and the imine isomer (C=N–C=O) corresponding to fructose is not produced due to a large energy disadvantage. Therefore, the dehydration of NAG provides a furan compound functionalised at the 3- and 5-positions (3A5AF), which is not observed in the glucose conversion. The dehydration of ADS requires a large amount of super-strong acid due to proton trapping by the amide group. We introduced the idea that H₃PO₃ can relieve this problem by producing esters with the substrate and easy protonation of the P=O moiety instead of OH groups. A new strategy like this is necessary to control the reactivity of NAG or its derivatives.

Regarding 3A5AF, we propose that the development of a Lewis acid catalyst is needed to improve the synthetic efficiency. Previous work used B(OH)₃ and Cl[−] with S/C ratios less than 1 to accomplish a good yield (up to 60%). Instead, we found that AlCl₃·6H₂O works for this reaction at a larger S/C (1.0) and a lower temperature (120 °C), which allowed us to increase the concentration of NAG up to 10 wt% and to isolate 3A5AF in a gram scale. However, the yield of 3A5AF was up to 30%. A new catalyst should be tolerant to amide and selectively activate OH groups for dehydration and H shift (Fig. 15), which further improves efficiency. After the synthesis of 3A5AF, its conversion to 3A5CF may open the door to bio-based polymers. The only method currently available is the iodoform reaction, and hence creating a new oxidation system is a fascinating subject.

We hope that the integration of these studies will achieve the practical utilisation of chitin toward establishing a sustainable carbon and nitrogen cycle.

Author contributions

HK: writing – original draft, visualisation. TS: writing – review & editing. AF: writing – review & editing.

Conflicts of interest

There are no conflicts to declare.

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