Ultrasound is defined as the science and technology of applying sound waves with frequencies above human hearing ability, essentially above 20 kHz [24]. Ultrasound can be classified into two main classes based upon frequency range i.e. power ultrasound and diagnostic ultrasound [25]. Power ultrasound is in the average range of 18-100 kHz and intensity >1 w/m² which can be applied in processing and chemical transformation, while diagnostic ultrasound with frequency above 500 MHz. Power ultrasound may also be referred to as Sonochemistry, which is the Ultrasound application in materials, chemical reaction and processes, excluding medical applications [26]. Another classification scheme categorizes ultrasound into three main classes based upon their frequency and power range as follows; low frequency high power ultrasound 20-10 kHz, medium frequency medium power ultrasound 100kHz-1MHz and the high frequency low power ultrasound >1 MHz.

Ultrasound with high impact applied in the food industry is in the range of low and high frequency. However, this definition is steadily changing with increasing and expanding applications of ultrasound in areas like cleaning, measuring quality and flow rate of foods [27,28]. This paper reviews the available literature on the application of power ultrasound to selected carbohydrates polymer and its sonochemo impact.

General Applications of Ultrasonic in Foods

Ultrasound is a form of energy carried by sound and can be transformed into other forms energy which is the basis of its applications [29,30]. Power ultrasound is capable of causing physical and chemical transformations through acoustic cavitation [31]. Ultrasonic treatment requires fluid for cavitation to be generated; moreover, some conditions must be met, for instance the frequency must be between 20 to 100 kHz.
Other conditions like solvent must meet required viscosity and polymer concentration should not be very high besides the solvent temperature should be low enough, approximately below 40°C [32].

Ultrasound is applied to food systems in the form of ultrasonic bath or probe [33]. Whereas in an ultrasonic bath, the solution of material to be sonicated is put in a container, thereafter the container is put in an ultrasonic bath with no direct contact between the material and equipment, in the ultrasonic probe, the probe is inserted in a container carrying the material allowing direct contact between the probe and the material [34]. The ultrasonic bath or probe creates cavitation in solution, owing to the formation of numerous water vapor bubbles which expand and contract rapidly, causing increased temperature and pressure up to 5000°C and 2000 bars respectively. Other forces which cause deformation is vibration, physical agitation and radical generation from the sonication fluid, water for instance is broken into H⁺ and OH⁻ ions which are radicals [35].

Ultrasonic machine

Ultrasonic machine consists of the following key parts:

Sonotrode: Sonotrodes are often referred to as the horn. They come in different sizes and power ratings. The tip transfers the oscillations into the medium to be sonified; eventually power is transferred through sonotrode into the material to be deformed [36].

Transducer: Ultrasonic transducer is the part that converts electrical energy into high frequency sound, the most commonly used one is piezoelectric transducers, even though in some cases magnetostrictive transducers are also used. High intensity low frequency ultrasound generates acoustic waves which are transferred in fluids as cavitation capable of causing chemical, physical or mechanical transformation in systems by creating micro cavities in liquids, numerous bubbles are formed and they collapse, the collapse rate is approximately 10-6sec causing intense heating and high pressure [37,38]. Second mechanism involves electronic waves causing increases in the amount of reactive species within the system [39].

Ultrasonic Power in Carbohydrate Polymer Degradation

Power ultrasound performance depends upon specific factors which influence cavitation. They include frequency, solvent, intensity and temperature [40,41].

Ultrasonic intensity

Ultrasound power is measured in watts; the absolute power equation is correlation between changes in temperature divided by change in time of a solvent system with specific heat capacity. The power equation was given by the following correlation:

Where P is absolute power (watts), “T” is temperature change, “t” is time difference, “Cp” is specific heat capacity of the solution and “M” is the mass of the solution.

The sonic intensity significantly contributes to ultrasonic effect in systems. For the probe type ultrasonic processor, the intensity of ultrasound power that is dissipated at the tip is predicted by the equation below [42].

Frequency

The frequency of ultrasound is believed to begin from 20 kHz, even though some authors state the minimum frequency is ≥ 16 kHz, the low frequency is able to generate cavitation’s, heating and surface instability in fluids. High power ultrasound is in the low frequency range between 20-100 kHz, while the high frequency ultrasound with frequency >500 MHz is applied in diagnostic work. Power ultrasound is able to cause physical, chemical and biochemical effects, which makes it applicable in Sonochemistry and processing.

Solvent

Action of ultrasound greatly depends upon solvent choice; the most commonly used solvent is water because most of the compounds are water soluble. However, some compounds are hydrophobic hence organic solvents are used in such cases. Solvents viscosity and surface tension inhibits sonication, therefore solvents with low surface tension and viscosity are preferred [43].

Temperature

Temperature is a crucial factor in several physical, biological and chemical reactions. Enzymatic and chemical reactions are known to be temperature dependent [44,45]. Ultrasonic is equally affected by temperature, with lower temperatures favors sonication, at higher temperature cavitation's collapse less violently, essentially high sonication occurs between 5°C-45°C [46]. Effective sonication depends upon balance between cavitation and temperature [47]. Where “I” is ultrasound intensity, “P” is input power and “r” is radius of the probe.

Mechanism of Polymer Sonication

Carbohydrate polymers consist of complex macromolecular structures ranging from starch, chitin, cellulose, pectin, carrageen, guar gum, alginate, dextrin and hemicelluloses etc. These polymers have little shared properties besides the basic Carbon, hydrogen and oxygen constituents [48]. Sonication of polymers is related to the polymer length, concentration and type of solvent, frequency of sound and intensity of ultrasound. So far Ultrasound has been applied in modification of several carbohydrate polymers including dextran, starch and chitosan [49], hydroxyl propyl methyl cellulose [50], carboxyl methyl cellulose, high methoxyl pectin [51], Guar gum and carrageenan gum [52,53].

While enzyme modification involves using different enzymes to break down specific bonds present in a polymer eg in pectin modification polygalacturonases are needed to depolymerize the Homogalacturonan chain, methylesterases are needed for demethylation and other enzymes to break specific bonds [54-57] similarly chemical method use specific chemicals which attack chemical bonds present in a polymer causing depolymerization [58]. These methods are laborious, expensive; require many cleaning steps and sometimes inefficient [59].

Generally, sonication breaks polymer chains at the center, which is the structurally weakest point. Molecules with high molecular weight and long chain length are more often broken at the center, than shorter and stiffer chains. Furthermore, the linear polymer chains are more easily sonolysed compared to branched ones. Therefore the initial molecular weight of the polymers plays an important role in the overall degradation rate. As polymers are broken down, a point is reached where the chains become so short and stiff, that further degradation is not possible and a lower molecular weight limit is attained [60]. The only way to overcome the lower limit is to increase the energy density of the sound field. It is worth noting that longer-chained molecules also lead to higher solution viscosity which also affects the rate of cavitation [61].
Sonication of pectin

Pectin modification consists of a process or step in which the pectin structure, physical and chemical property is changed to suit a given application. Methods like chemical [62,63], heat modification [64], irradiation [65] and enzymatic [66] have been applied in pectin modification. Sonication has been applied in pectin to a lesser extent with only handful of published papers to that effect [67,68].

Mechanisms: Sonication of citrus pectin was reported and observed that the average molecular weight decreased from 464 to 296 k Da after 30 minutes of treatment, degree of methylation reduced slightly, the neutral sugar side chain was degraded while the total monosaccharide content remained the same. The rate of pectin molecular weight degradation was predicted by the following equation

\[ \frac{1}{M_t} - \frac{1}{M_0} = kt \]  

(1)

Where k is the rate constant (mol/g·min) of molecular weight during sonochemical degradation, where "t" is the sonochemical degradation time (min); "Mt" and "M0" are the weight-average molar mass at time t and at time 0 (Daltons), respectively.

While sonicating citrus pectin Liu et al., noted decrease in the molecular weight of sonicated citrus pectin reduced from 630 k Da to 230 k Da, esterification degree and viscosity with increasing time from 0 to 90 min, ultrasound intensity and ultrasound duty cycle. Optimum sonolysis was noted at sonication intensity 302 W/cm², above this intensity the sonolysis activity showed no significant change.

In a study on the effect of high-intensity ultrasound on the rheological and optical properties of high-esterified apple pectin dispersions, resulted in decreased turbidity, weaker gel and decreased in viscosity, which were attributed to decreased molecular weight at the highest intensity level of 40 W cm⁻² in 30 min. This is similar to other authors who showed decreasing molecular weight, viscosity and gel strength during sonication [69,70].

Additionally, Sonication was applied in preparation of pectosomes and chitosomes for liposomes drug delivery system. The presence of pectin and chitin stabilized the retention of metronidazole; moreover, sonication of the liposomes resulted into nano-sized particles with size ranging from 62 nm-222 nm for freshly prepared liposomes after sonication.

Characteristics of sonicated pectin: The sonicated pectin products have been characterized by few authors. Some noted changes included reduced degree of polymerization, molecular weight, reduced DM and neutral sugar degradation. Sonication can be used to make micro and nano-sized pectin derivatives for different applications.

Potential applications of modified pectin: Pectinosomes have been shown to be a good candidate for metronidazole delivery system, owing to their mucoadhesive properties enabling them to have a longer residence time in the vagina. Modified pectin from various processes like pH and heat treated one were tested for its anticancer activity and it showed higher potency than normal pectin. The most well-known pectin is modified citrus pectin (MCP), which is prepared using inorganic acid (HCl) and alkali (NaOH), which lead to de-esterification of pectin and de-esterification of homogalacatan [71]. Similar changes were observed when enzymatic and chemical methods were used to modify pectin [72,73].

Modified pectins anticancer activity is linked to induction of apoptosis and reducing proliferation of cancer cells [74-77]. MCP is commercially marketed as a nutraceutical and chemotherapeutic substance against several cancers [78]. In another study modified sweet potato pectin was shown to possess capacity to reduce cancer cell proliferation [79]. Low Degree of Polymerization (LDP) pectin have been shown to be more potent compared to high DP ones partly owing to their ability to be transported across intestinal wall and into the blood stream. Finally, oligo pectins are capable of inducing plant immunity, thus enabling plants to boost their immune system against pest attack. Prebiotics, oligo pectins have been shown to be better prebiotics for specific probiotic bacteria like Bifidobacteria and Lactobacilli. The selectivity of oligosaccharide depend on the chain length and presence of specific functional groups, the oligo pectins with short chain being preferred substrate for probiotics [80,81].

Carrageenan

Carrageenan, a sulphated polysaccharide extracted from certain species of red seaweed [82]. Carrageenan exist in different forms of κ-kappa with single sulphate group, i -iota with two sulphate groups and λ-lambda with three sulphate groups. Carrageenan form thermo-reversible gels with potassium ions or other metallic ions, it is also capable of forming gels with proteins hence often applied in chocolate milk and fresh cheese. Other applications include processed meats, dietetic formulations, infant formula, toothpaste, cosmetics, skin preparations and laxatives.

Sonication of carrageenan: A recently sonication of carrageenan solution was investigated at three frequencies of 23, 48 and 83 kHz at intensity of 50 W. The group observed higher reduction in shear viscosity at 45 kHz with full shear viscosity recovery on stopping ultrasound application, the other frequencies 23 and 83 kHz had low effect, probably due to their inability to build effective bubbles and create cavitation due to low frequency at 23 kHz and too high frequency at 83kHz respectively. During sonication free OH-ions build up in the system, this is attributed to splitting of water molecules into hydroxyl and hydroxyl radical species.

In an earlier study carrageenan was degraded by sonication, the rate of degradation increased with increased intensity, time, and reduced pH and concentration [83]. K-carrageenan had a higher degradation rate compared to ξ-carrageenan; it is probably because the presence of additional sulphate groups in ξ-carrageenan could have contributed to its less vulnerability to sonication.

Possible application of carrageenan oligomers: Sonication offers an effective method of carrageenan oligomer processing. The carrageenan oligomers can be applied as plant growth promoter, antiviral, antitumor, antioxidants, anticoagulant, antithrombotic and hydrogels for burns dressings. Each function is linked to oligomers with specific molecular weight range [84].

Guar gum

Guar gum Guar gum is obtained from endosperm of the guar plant, Cyamopsis Tetragonolobus. Guar gum is a galactomannan with a linear backbone of β-D-(1, 4)-mannose irregularly substituted by uncharged α-D-(1, 6)-linked galactose side groups [85]. It is cold water soluble producing highly viscous pseudo plastic solution which makes it preferable choice in industry. Guar gum is industrially used as gelling, viscosifying and thickening agent. The viscosity of guar depends on the temperature, pH, time, concentrations, degree of agitation, and particle size of the gum [86].
Sonication of guar gum: The only available study documented on sonication of guar gum is reported by Ansari et al., which revealed that sonication could also be a convenient method of reducing the molecular weight of Guar Gum. Furthermore, the NMR results demonstrated that sonication did not alter the structural characteristics of the polymeric chain of guar gum. The depolymerization of Guar gum by sonication resulted in lower molecular weight fractions within three minutes of sonication the MW of initial GM decreased by half. Difference in water uptake and anisotropic elongation is also a significant achievement of sonication.

Applications of modified guar gum: The reduction of Guar Gum (GG) molecular weight for an appropriate modulation of its flow and gelling properties could ultimately be useful in pharmaceutical industries for innovative drug formulations, it could as well be applied in cosmetics and food industry.

Chitin and chitosan

Chitin a biopolymer consisting of poly(β-(1→4)-N-acetyl-D-glucosamine), it is a natural polysaccharide derived from several living organisms; chitin is the second most abundant polymer after cellulose in the world. Chitosan is the most significant derivative of Chitin obtained after partial deacetylation. Chitosan is a linear polysaccharide composed of β-(1→4)-linked 2-amino-2-deoxy-D-glucopyranose (GlcN, D-unit) and 2-acetamido-2-deoxy-D-glucopyranose (GlcNAc, A-unit) [87].

Sonication of chitin: Previously chitosan was prepared using chemical and enzymatic methods [88,89]. Investigation on impact of ultrasound on chitosan molecular weight and degree acetylation revealed that 1% Chitosan in 1% acetic acid solution was sonicated at 16.5, 28.0 and 35.2Wcm⁻² for 0-30 minutes, the researchers noted that the degree of acetylation was not changed while the molecular weight was reduced, the reduced sonication rate after optimum molecular weight reduction rate was attributed to many molecules with more bonds to be broken hence the rate of reducing molecular rate reduced with time after a given critical time limit. Finally, the group concluded that the effect of ultrasound was majorly affected by sonication time and power [90].

In another research to investigate the involvement of mechano-chemistry and radical reaction during sonication to reduce chitosan molecular weight by employing ultrasound of 230 kHz frequency and treatment time 0-90 minutes. Hydroxyl radicals contributed to approximately 60% of molecular weight reduction, the average sonolysis rate was 8×10⁻¹ⁱ mol J⁻¹. Optimal molecular weight reduction was noted at 80 minutes and continuous treatment did not lead to further molecular weight reduction, this was attributed to increased number of molecules to be broken. The results further showed that degree of deacetylation of chitosan was not altered at all the power and time.

Moreover, Liu and co-workers sonicated shrimp shell chitosan at 800C, 250 W and 0-99 h. The initial chitosan fractions had the following molecular weight 65.5×10⁴ K Da, 154×10⁴ K Da, 77.5×10⁴ K Da and 44.7×10⁴ K Da and with de-acetylation degree DD% 61.9, 72.1, 87.1, 91.6 respectively. Results showed that chitosan with higher molecular weight and higher de-acetylation degree was more easily depolymerized, The DD degradation was inversely correlated, with the chitosan having DD >90% having no detectable degradation while ones with DD <80% showed significant degradation. Besides, polydispersity decreased with treatment time [91].

Applications of chitosan: Chitosan is widely used in a range of diverse fields, including waste management, medicine, food and agriculture. Chitosan has biological properties such as biocompatibility, antimicrobial, biodegradability, mucoadhesion, anticholesterolemic and permeation enhancement effects [92,93]. The molecular weight of chitosan is a crucial factor in its functionality. Low molecular weight chitosan have been shown to possess much higher bioactivity than the large molecular weight ones. Some of the possible applications of sonicated chitosan are as follows:

Chitosan as antimicrobial agents: Ultrasonically treated chitosan showed a higher antimicrobial activity against known food pathogens; E. coli and S. aureus. The increased activity was attributed mainly to reduced molecular weight, particularly chitosan of molecular weight range of 5-10 K Da. The use of chitosan in bioactive packaging material against Pseudomonas aeruginosa and Staphylococcus aureus was explored; it showed effective anti-microbial activity [94].

Anti-diabetes mellitus: Low molecular weight chitosan with molecular weight of 20 kDa has shown to lower the progression of diabetes mellitus. Other possible function of chitosan and oligochitosans include hypcholesterolemic, immune-stimulating, antitumor and anticanicancer effects. It also exhibits accelerating calcium and iron absorption, anti-inflammatory, antioxidant and Angiotensin-I-converting enzyme (ACE) inhibitory activity-regulating blood pressure.

Food applications of chitosan: Chitosan is widely used in food industries as dietary fiber, lipids binding agents, preservative, thickener and stabilizer. It is also use as protective fungi static, antibacterial coating for fruit. Low molecular weight chitosan is used as nutraceuticals, antimycotic and antimicrobial agents. Moreover, chitosan carrageenan nano fibrils have shown a good potential to be used as nano composite packaging material [95].

Other biomedical applications: Chitosan has various biomedical applications like surgical sutures, dental implants, artificial skin, rebuilding of bone, corneal contact lenses, time release drugs for animals and humans encapsulating material. Besides, it equally possesses Immunological, mucoadhesive, antitumor hemostatic and anticoagulant, healing, bacteriostatic ability. Chitosomes are also an excellent drug loading and delivering system [96].

Starch

Starch is of the most abundant natural polymer. It consists of amylose and amylopectin, starch is a partially crystalline polymeric substance. X-ray diffraction patterns of native starch granules may give A, B, C and V-type depending on the source of starch [97]. The molecular weight of starch is variable depending on its source. Starches and modified starches can be used to affect the physical properties of many foods [98].

Sonication of starch: Starch modification is achieved through different methods [99]. Acid and enzymatic modification of starch has been used to make nano-colloid or crystalline starch. It has been reported that ultrasonic treatment also reduced molecular weight of corn starch; the ability of ultrasound to degrade starch at 360 kHz was shown to be mediated by OH- and mechano-chemical effects. Starch had a lower yield rate compared to chitosan because starch is highly branched while chitosan is linear. It was observed that chain breakage was directly related to time up to around 15 minutes thereafter the rate experienced slight decline.

In another study on the effect of ultrasonic on physical properties of corn starch, it was found that the gelatinization temperature was not
changed. The rheological property of ultrasonic treated starch showed a reduced consistency coefficient. However, it is worth noting that the treatment was only done at two time points of 15 and 30 minutes, while in the previous studies impact of ultrasound was high after longer time treatment.

Effect of ultrasound on viscosity of various starches was investigated, sonication leads to marked reduced viscosity of waxy maize, tapioca, potato and sweet potato starches. Sonication decreased starch molecular size by de-polymerization, the rate of de-polymerization was high up to 30 minutes then it reduced gradually with time and nearly levels off after 60 minutes [100].

The first case of making nano starch particles using purely physical treatment was done by Bel Haaj and coworkers, the ultrasound was applied in waxy corn and standard starch, they observed conversion of starch into Nano sized particles with size range 100-200 nm using high intensity and long treatment time, moreover, much smaller nanoparticles less than 100 nm were generated with sonication for 60 minutes yielding 80 nm and 90 minutes. Moreover, the starch granular structure was destroyed after 75 minutes of sonication to 30 nm.

Applications of low degree of polymerization (DP) starch: Sonicated starch generally has low degree of polymerization owing to depolymerization. Nano scale starch has been made by ultrasonic treatment; this creates materials for nano-starch applications like improved packaging materials. Nano composites can be used as fillers materials for biodegradable materials and packages. Additionally, Nano-starch improves the properties of packaging materials, it can also improve the structure and texture of foods where applied.

Future Prospects of Ultrasonic in Polysaccharide

The previous research results have shown not only the promising prospect of using ultrasound in polymer modification, but also in extraction of compounds [101] and processing [102,103]. Industries are always looking for innovative, affordable, effective and efficient methods of processing, modification and extraction of compounds [104-106]. Food and pharmaceutical ingredients industry can easily bank on the process. Food and pharmaceutical ingredients industry can easily bank on the process.

Polysaccharides being the most commonly used food ingredients but suffering from limited physico-chemical characteristics which limit their application, the need to modify is pressing. The growth of functional foods, pharmaceutical and cosmetic is driving sonication of polysaccharides since sonication is more adaptable, efficient and effective in several processes, it is mostly likely going to become one of the main technologies that will transform industries in the coming years. In summary, sonication is an effective and efficient method for polysaccharide modification and future oligosaccharide and nano scale molecules processing would mostly probably be done by sonication process. The sonolysis of the reviewed polysaccharides could be used to predict to some extent its activity in other polysaccharides like xanthan, alginate, carboxymethyl cellulose, konjac gum, gum arabica etc. Future study should be targeted at optimizing the sonication parameters, deeper understanding of mechanism, exploring sonicated products application and up-scaling the process.

Conclusions

The future of sonochemistry is getting brighter with increasing research endeavors, good research results and up scaling in the discipline. Sonochemistry is a green technology which is cheaper, more adaptable, efficient and effective in several processes, it is likely going to become one of the main technologies that will transform industries in the coming years. In summary, sonochemistry is an effective and efficient method for polysaccharide modification and future oligosaccharide and nano scale molecules processing would mostly probably be done by sonication process. The sonolysis of the reviewed polysaccharides could be used to predict to some extent its activity in other polysaccharides like xanthan, alginate, carboxymethyl cellulose, konjac gum, gum arabica etc. Future study should be targeted at optimizing the sonication parameters, deeper understanding of mechanism, exploring sonicated products application and up-scaling the process.

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