### **Green Chemistry**



### **CRITICAL REVIEW**

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# Biomass-derived functional materials as carriers for enzymes: towards sustainable and robust biocatalysts

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The unique catalytic properties of enzymes have led to the production of useful medicinal intermediates, foods, and biofuels from sustainable sources. However, the instability of soluble/free enzymes under several challenging conditions (e.g., pH, proteolysis, temperature, ionic potential, chemical denaturants) restricts the use of enzyme-based biocatalysts. Encapsulation of enzymes on suitable carriers would mitigate the instability issues faced in robust biocatalysis. An "ideal" carrier material employed for protein immobilization should be nontoxic, scalable, biocompatible, and should not compromise the biological activity and structure of proteins/enzymes. Thus, biodegradable and renewable biomass-derived functional materials (BDFMs) are envisaged as promising carriers for enzymes. BDFMs have in-built chemical functionalities and desirable physicochemical properties that enable their use in enzyme catalysis at the industrial scale. Numerous BDFMs have been used as immobilization matrices to improve the biocatalytic activity and stability of various enzymes. These solid materials are renewable and environmentally friendly compared with synthetic polymers. This review highlights the advancements, challenges and prospects in the emerging field of BDFMs (cellulose, silk protein, chitin, chitosan, lignocellulose, and a combination of biopolymers such as chitin/lignin and chitosan/alginate) for immobilization of enzymes (e.g., α-chymotrypsin, cytochrome c, carbonic anhydrase, glucose oxidase, ribonuclease, cholesterol oxidase, alkaline phosphatase, β-glucosidase, lipase, horseradish peroxidase, catalase, tyrosinase, acetylcholinesterase, amylase, invertase, protease, laccase, β-galactosidase, and several others) for biocatalytic processes. This review also describes the relationship between the structural properties and functionality of several enzymes immobilized in BDFMs, and profiles the impact of pH, temperature, reusability, stability of storage, and the activity of these enzymes. Future perspectives in this promising field, as well as potential difficulties, are discussed. This review will help in refining biocatalysis technologies whereby biomassderived, environmentally friendly materials are employed as enzyme supports.

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#### 1. Introduction

Society has high dependence on non-renewable resources, which is unsustainable in the long run. Increases in population sizes and environmental pollution have prompted strategies for "green" chemistry, sustainable resources for fuels, chemicals, and materials, and reducing waste. <sup>1,2</sup> In this scenario, the development of flexible and integrated "biorefineries"

to produce biofuels and bioproducts from renewable biomass sources can aid transition to a novel "bioeconomy" for more efficient and sustainable global development. The "circular economy" concept is taking a central role in sustainable development and, for this reason, deserves attention.<sup>3-6</sup> Renewable denotes sustainable and abundantly available feedstocks for the production of biofuel and biochemicals via suitable bioconversion. "Biocatalysis" is a charming technology used in various industrial applications. An enzyme or biocatalyst is essentially a non-hazardous, non-toxic material that is easily obtained from widely available renewable resources. Enzymes are Nature's sustainable catalysts which drive a variety of reactions. In accordance with the theory of green chemistry, biocatalytic processes are more sustainable, environmentally friendly, and cost-effective than traditional chemical processes. In the context of green chemistry and sustainable development, the utilization of enzymes as efficient catalysts offers

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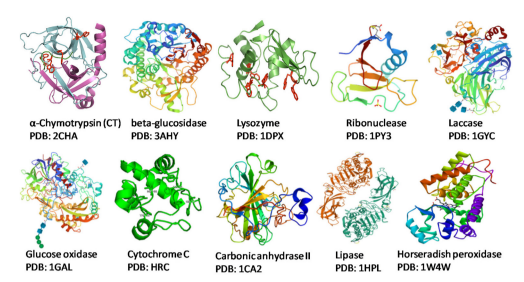


Fig. 1 Structures of commonly used enzymes. Source: Protein Data Bank.

several interesting characteristics.<sup>1,2</sup> Enzymes are biodegradable, reusable biocatalysts which do not create by-products. Enzymes are made up of several amino acids that are linked together by a peptide bond (-CONH-).8,9 The native state of a protein is a result of complex interactions (e.g., hydrogen bonds, van der Waals interactions, ionic interactions, hydrophobic interactions) which ultimately provide stability to proteins and prevent ruinous conformational changes.<sup>10</sup> Each enzyme exhibits a great degree of chemo-, regio- and stereospecificity towards a substrate because of the binding properties at the active site. The native or tertiary structure of some common enzymes that have been modified with biomass-derived functional materials (BDFMs) is shown in Fig. 1.

In the synthesis of complex pharmaceuticals, enzymatic biocatalytic approaches are more environmentally friendly and sustainable than chemical methods. Due to their capacity to catalyse reactions with high efficiency and specificity, they have emerged as preferred tools in green chemistry and are being used more frequently in industrial processes. 11 Fig. 2 shows various factors that have an impact on the stability and activity of enzymes. In recent years, biocatalysis has emerged as an imperative technology for meeting the growing demands for green and sustainable chemical manufacturing. However, enzymes are moderately stable in terms of their structural and chemical stabilities, and enzymatic processes are conducted under physiological conditions in a buffer, with high rates and selectivities. Harsh processing conditions, such as the presence



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protein stabilization using a wide variety of "green" solvents (e.g., ionic liquids and deep eutectic solvents). She has developed a green, innovative, and cost-effective technology for the enhanced dissolution and extraction of bioactive compounds from sustainable biomass.



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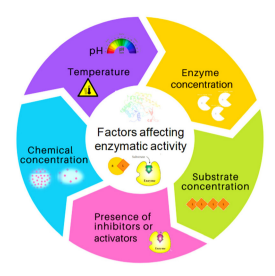


Fig. 2 Various factors that influence the activity and stability of enzymes/proteins.

of volatile organic solvents, extreme pH, and temperature, are the main barriers to the effective use of enzymes in therapeutics and as biocatalysts. Moreover, a lack of long-term activity, stability, reusability, and a challenging recovery rate can hinder the industrial application of enzymes.<sup>1,2</sup> To overcome these problems, numerous solutions have been postulated by researchers, such as methods for protein engineering, chemical modifications, excipient addition, and immobilization.12

The progress of effective immobilization strategies has paved the way for enhancing the recovery, recycling, operational stability, and storage of enzymes.<sup>13</sup> Immobilization is also very important for effective biocatalysis. Enzyme immobilization is the stable fixation of the protein within a solid support/matrix by physical or chemical interactions (or both)

so that the enzyme remains fully functional (or at least retains most of its catalytic activity). There are various methods for enzyme immobilization: adsorption, entrapment, covalent binding, and crosslinking. 13-16 During immobilization, the integrity of the structure and the function of an enzyme are at great risk, and the biocatalytic performance may be compromised. In addition, the co-immobilization of two or more enzymes can afford multifunctional solid biocatalysts capable of catalysing processes in biocatalytic cascades. 14 Therefore, immobilization of enzymes is an essential method that has a significant effect on the stability and effectiveness of enzymes. Several strategies for enzyme immobilization have been categorized: adsorption binding, entrapment, covalent binding, and crosslinking with/without a support. 15,16 Immobilized proteins have been widely use in industrial applications (e.g., analytical, pharmaceutical, commodity chemicals, food, and cosmetic industries) as well as energy production and biomedicine. 15,16 Therefore, immobilization technologies must be improved and diversified to support the development formats, better economies, and higher performance. 13,14 Supports can be made of hybrid materials, inorganic materials, metal-organic frameworks (MOFs), nanoflowers, hydrogels, polymers, or biomass-based materials. 15,17,18-21 BDFMs are exceptional supports for immobilising enzymes to enhance the activity and selectivity of enzymes.21,22 BDFMs have attracted much interest due to their abundance, environmental friendliness, sustainability, and unique structures.1 Biomass-based matrices have great microbiological resistance, biocompatibility, robust mechanical properties, acceptable stability, and controllable biodegradability, which might offer a favourable microenvironment for enzyme stability. BDFMs with superior properties and precise morphology for improving facile biocatalysis is a research "hotspot" (Fig. 3). BDFMs are created by transforming the



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materials science. Understanding the fundamental mechanisms of green chemistry and the interaction between plants and nanomaterials is the main focus of his research team.

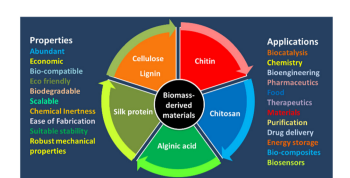


Fig. 3 Potential applications of BDFMs in biocatalytic processes.

natural resources (plants, food, animals, microorganisms) and their wastes into high-performance materials using physical or chemical processes.<sup>2</sup> BDFMs promote the stabilization of entrained molecules, such as therapeutic proteins or enzymes, without compromising their activity.<sup>23</sup> Furthermore, because BDFMs have been approved by the US Food and Drug Administration, these supports/matrices are ingestible and may have useful applications *in vivo*.<sup>23</sup> The immobilization of industrial enzymes would enable the reuse of expensive enzymes for enhanced utility in industrial processes, whereas the immobilization of bioactive enzymes might be incorporated into diagnostic and therapeutic applications.<sup>24</sup> In this review, we evaluate available data regarding the potential of BDFMs as sustainable carriers for enzyme stabilization.

Precursors for BDFMs are readily available, inexpensive, and primarily produced from plants (cellulose, lignin, alginate), animals (silk, chitin, chitosan), and microorganisms (bacterial cellulose). BDFMs possess numerous desirable inherent qualities that point to their potential use as an



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enzyme-stabilisation matrices to protect them from different stresses. This review focuses on the main categories of biomass-based support materials for different enzymes, or biocatalyst designs/functionalization for improving the stability and activity of enzymes and maximizing reaction efficiencies.

Fig. 4 depicts an overlay analysis which focuses on evaluating the qualities and academic importance describing BDFM use to increase enzymatic efficiency. An extensive evaluation of the academic and scientific literature accessible in this field of research was carried out using the Web of Science (WOS). In the current review, VOS viewer was used to display the cooccurrence of keywords, and an overlay map was developed. The circles in the overlay map depict the occurrence of the keywords in the chosen dataset (Fig. 4). The keyword co-occurrence map was created by choosing keywords that appeared ≥10 times. As a result, 315 out of 7211 keywords met the criteria and were further classified into eight major clusters. As shown in Fig. 4, since 2000, there have been sporadic reports on enzyme immobilizations using biomass-derived materials. In the years between 2008 and 2018, biomass-related research has experienced explosive growth. The important discoveries made possible by the use of biomass-derived materials as carriers for long-term stability and enhancing the catalytic activity of enzymes and proteins are summarised in this overview. Recent biocatalytic contributions of biomass-based supports are discussed in the next sections.

### 2. Lignocellulosic-based supports (carriers) for enzyme immobilization

Lignocellulose and cellulose-based materials are attractive support materials for enzyme immobilization because they have high porosity, strong mechanical properties, are highly resistant to microorganisms, as well as having tunable biodegradability, excellent stability, and dense network of hydrogen bond crosslinks. 25-27 Being the most abundant polysaccharide, cellulose is made up of linear chains of p-glucose units connected through  $\beta(1 \rightarrow 4)$ -fashioned glycosidic linkages, as well as intra and intermolecular H-bonding (Fig. 5). This results in the formation of cellulose microfibrils that are firm and strong. To create  $\beta(1 \rightarrow 4)$  glycosidic bonds, every alternate glucose molecule in cellulose is inverted. Cellulose-functionalized materials can be made from cellulosic sources such as cotton and bacterial cellulose wherein the hydrolysis of cellulose generates nano-sized crystalline material.26-28 Several such forms of cellulose, such as cellulose hydrogel microspheres (CHMs), cellulose nanocrystal (CNCs), cellulose nanowhiskers (CNWs), and nanofibrillated cellulose (NFC) have been studied as possible supports for enzyme immobilization. They are characterized by high enzyme loading, enhanced stability, improved activity, and increased mass transfer rate. 29-47 Various immobilization techniques can be applied to load enzymes onto the matrices of lignocellulosic biomass/ modified cellulose-based materials for the conjugation and stabilization of enzymes and proteins. Recently, Ren et al. 45

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microbial cellulose biomaterials @ in-vitro chemical-modification drug-delivery lysozyme activation polymerization bacterial cellulose biocompatibility penicillin-g acylase oligosaccharide chemistry invertase glutaraldehyde nanocomposites membranes supports chitin electron=transfer enzyme immobilization fabrication glucoamylase thermostability biocatalyst enzymes nanoparticles silk fibroin electrochemistry biodiesel horseradish-peroxidase xylanase performance reusability hydrogen peroxide biosensor Jaccase bioconversion regenerated silk fibroin beta-glucosidase glucose

oxidation

ionic liquid

biosynthesis /

enhancement

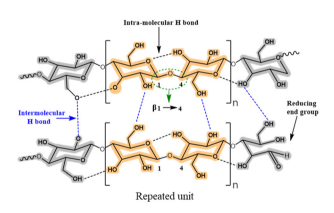
trametes-versicolor

Fig. 4 Overlay map of co-occurrence of keywords concerning BDFMs as carriers for enzymes.

biofuels

detoxification

alginate beads



ultrafiltration

fermentation

lignocellulose

VOSviewer

pretreatment biomass

bioethanol enzymatic-hydrolysis

Fig. 5 Intra- and inter-molecular hydrogen bonding and possible reaction sites in cellulose.

reported the superior potential of delignified bamboo over the immobilization of β-glucuronidase (BGU) with a significantly higher loading capacity (~8 mg m<sup>-2</sup>·g<sup>-1</sup>), recyclability (13 cycles), and longer shelf life (≤7 weeks). Cellulose matrices were employed to enhance the robustness of the enzyme in analytical and synthetic applications for hierarchical flowbased bioreactors.

Uth and co-workers<sup>47</sup> used a modular approach for sitedirected, biorthogonal protein immobilization. A general twostep method was created that took advantage of extremely effective oxime ligation along with enzyme-mediated protein

coupling onto the surface of peptide-modified crystalline nanocellulose. Conjugate analyses revealed that, after being grafted onto CNC, the immobilized glucose oxidase (GOx) maintained its activity (Fig. 6). To improve the salt and thermal tolerance of cellulase enzyme, Montamedi and colleagues<sup>46</sup> immobilized its cloned form (PersiCeL3) over a carboxymethyl cellulose (CMC) hydrogel matrix. With halotolerant behaviour >65% and almost two-folds higher activity than the native enzyme, immobilized PersiCel3 showed superb performance which could be used in lignocellulosic biomass industries to boost the production of value-added products in hot and salty environments.

ctrodeposition

2020

polypyrrole

2010

amperometry

tyrosinase

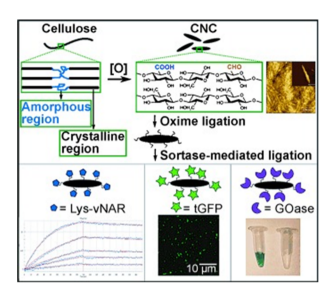
pesticides

2000

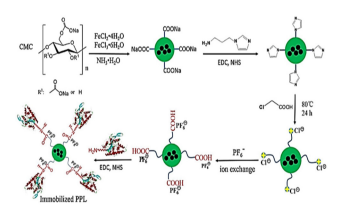
construction

Authors transformed the primary alcohols on the surface of CNC to the corresponding aldehyde functionality, which enabled immobilization of bioactive modules via extremely effective oxime ligation. Immobilization of the enzyme on the CMC-hydrogel matrix could be used to improve the robustness of enzymes with strong resistance towards high temperatures and heavy salt concentrations.7,46 Suo et al.7 prepared ionic liquid (IL)-modified magnetic CMC nanoparticles on which porcine pancreatic lipase (PPL) was immobilized covalently (Fig. 7). The immobilized lipase (PPL-IL-MCMC) demonstrated specific activities that were 1.43- and 2.81-fold greater than those of free PPL and PPL-MCMC, respectively, with simple and efficient recoverability.

Extending the work further, the authors studied the effect of urea (Fig. 8a) and thermal stresses (Fig. 8b) over the activity

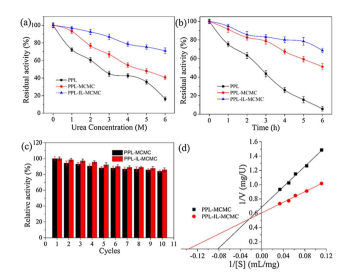


**Fig. 6** Postulated chemoenzymatic approach to protein immobilization onto crystalline cellulose nanoscaffolds (schematic). Reproduced from ref. 47 with permission from Wiley-VCH, copyright 2014.



**Fig. 7** A common method for preparation of IL-modified BDFMs and enzyme immobilization upon it. Reproduced from ref. 7 with permission from Elsevier, copyright 2020.

and stability of PPL. The conjugate retained >68% activity but the native enzyme could barely maintain its functions. The immobilized enzymes PPL-MCMC and PLL-IL-MCMC could maintain 40.8% and 70.9% of their original activity, respectively, at a urea concentration of 6 mol L<sup>-1</sup>, whereas the native enzymes retained only 15% of their activity. A similar trend was reported for thermal stress for a long time (Fig. 8b). The advantage of combining two stabilizing systems can be clearly envisioned by recyclability and Michaelis-Menten parameters (Fig. 8c and d). This is due to ion interactions, covalent bonds, H-bonds, and  $\pi$ - $\pi$  stacking between the carrier and enzyme. This could also support maintenance of the stiffness and integrity of the enzyme structure under harsh conditions, thereby stopping the enzymes from becoming partially inactivated.7 Utilizing halloysite nanotubes (HNTs) as a template, Sillu et al., 48 described the synthesis of a nanobiocatalyst that



**Fig. 8** Effect of the CMC-hydrogel matrix on the stability (a–c) and kinetic parameters (d) of lipases. Reproduced from ref. 7 with permission from Elsevier, copyright 2020.

immobilized the enzyme that catalysed the breakdown of cellulose into glucose: cellulase.

As compared with free enzyme, immobilized cellulase showed greater stability at high temperatures (≥60 °C) and storage capacity and activity. Luo and co-workers<sup>49</sup> created porous magnetic cellulose microspheres (MCMs) activated by epoxy chloropropane to boost the covalent immobilization of penicillin G acylase (PGA) to increase the effectiveness of catalyst recovery. The immobilized PGA displayed strong catalytic activity, improved pH tolerance, and enhanced thermal stability. The catalysts were recovered by separating the enzymeloaded matrix from the reaction solution. In the quest of producing protein-friendly biomaterial for biocatalysis and biotechnological applications, our research team<sup>50</sup> synthesized tendril-like functional carbon helices (TLFCHs) from lignocellulosic biomass via a "green" solvothermal technique utilising deep eutectic solvent (DES) as a soft template and catalyst. Exploiting the benefits of helicity and in-built chemical functionalities, we immobilized cytochrome c (Cyt c) on the surface of TLFCHs (Fig. 9). At optimized conditions, the specific activity, pH stability, and peroxidase activity of Cyt c were increased without impacting the structural integrity of the protein. Furthermore, lignocellulose-based hydrogels possess the advantages of low cost, biocompatibility, biodegradability, hydrophilicity, non-toxicity and, most importantly, controllable properties. 17,51-62 Hydrogels based on lignocellulose have revealed promising applications in biocatalytic, biomedical, and bioelectronic areas. 51,52 Jo et al. 51 generated cellulose hydrogel microspheres by sol-gel transition employing a 1-ethyl-3-methylimidazolium acetate ([Emim][Ac])-in-oil emulsion for immobilization of lipase. Immobilized lipase showed greater efficiency as compared with the native enzyme. In contrast to lipase immobilized on MCC or mm-sized hydrogel beads, lipase immobilized on cellulose microspheres demon-

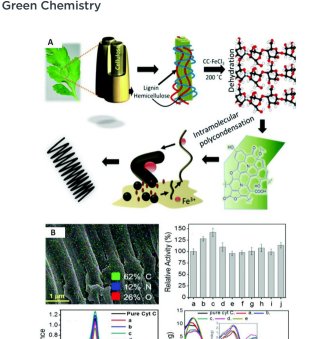


Fig. 9 Schematic depictions of (A) a plausible mechanism of the sequential growth of TLFCHs from *Parthenium* biomass during a solvothermal process in the presence of a DES (CC: FeCl<sub>3</sub>, 1:2 mol ratio) (B) Cyt C immobilized on TLECH and showing activity and stability results (ref. 50).

700

210 220 230

Wavelength(nm)

500

strated a much greater loading efficiency, immobilization yield, and specificity constant.<sup>51</sup> In another study,<sup>52</sup> cellulose and lignin were co-dissolved in an IL to produce cellulose/ lignin composite hydrogel beads. The activity, protein loading, and specific activity of the lipase immobilized on the cellulose/lignin beads were 2.6-, 2.2-, and 1.2-times higher than those of the lipase immobilized on cellulose beads, respectively. 52 Kim et al. 53 extended this work by conjugating cellulose with other bio-polymers, such as chitosan, carrageenan, agarose and agar. Candida rugose lipase (CRL) immobilized over these hydrogels in the presence of [Emim][Ac] demonstrated greater immobilization yield compared with that of cellulose beads.<sup>53</sup> In addition to the cellulose materials mentioned above, chemically altered nanoscale cellulose materials make excellent matrices for enzyme immobilization. Arola and co-workers<sup>17</sup> immobilized proteins over NFC through amine, epoxy, and carboxylic-acid functionalization. 17 The NFC structure is beneficial for the stability and catalytic activity of proteins.17 Organophosphorus hydrolase (OPH) Flavobacterium ATCC 27551 was immobilized on powdered plant cellulose treated with epoxy (Fig. 10) by Sharifi et al.54 The immobilized OPH demonstrated better storage, temperature, pH, and reusability properties than the free enzyme. Apart from epoxy groups, introducing aldehyde and carboxyl groups on cellulose matrices is an effective approach to attach them with the amino groups of enzymes. 56,57

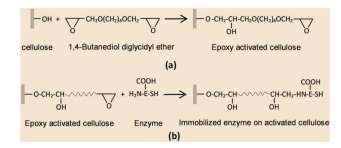


Fig. 10 Plausible reaction mechanism of enzyme immobilization through the epoxy method. (a) Surface reaction of a cellulose matrix with 1,4-butanediol diglycidyl ether (BTDE). (b) Covalent coupling of enzyme onto the surface of activated cellulose. Reproduced from ref. 54 with permission from Elsevier, copyright 2018.

Different chemical processes are used to oxidise the hydroxyl groups in cellulose to produce aldehyde or carboxyl groups. For example, Hao  $et~al.^{56}$  used sodium periodate (NaIO<sub>4</sub>) for oxidizing cotton fibres to develop aldehyde groups in them. <sup>56</sup> A similar study was reported for oxidation of cotton yarn by NaIO<sub>4</sub> to introduce aldehyde groups and further used for immobilization of trypsin. <sup>57</sup> The highest concentration of immobilized trypsin was 6.1 mg g<sup>-1</sup> of dry cotton yarn. Over 60 days of storage, the activity of immobilized trypsin increased, and revealed >90% and >70% of the initial activity at 4 °C and 25 °C, respectively. <sup>57</sup>

In another study by the Gong group,<sup>58</sup> NaIO<sub>4</sub> first oxidised the cellulose in a loofah sponge to produce aldehyde groups. Then, the oxidised loofah sponge was used as a carrier for the covalent immobilization of lipase. The immobilized lipase provided better thermal stability, storage stability, and reusability. Besides periodate, 2,2,6,6-tetramethylpiperidine-1-oxyl radical (TEMPO) was used by Qian and co-authors to control the distribution of carboxylic groups over CNF macrogels, over which phospholipase from *Thermomyces lanuginosus* was immobilized and showed great reusability.<sup>43</sup>

To cutdown the time frame of purification and immobilization processes, Gennari et al., 59 engineered a single-step methodology by altering the surface of MCC with CBD. β-Galactosidase from Kluveromyces sp. was simultaneously purified and immobilized over this carrier and showed high stability at up to 40 cycles for the hydrolysis of milk lactose. Various other types of cellulose matrices have been used as carriers for enzyme immobilization (Table 1).63-85 To improve the tolerability of enzymes in enzymatic membrane bioreactors (EMBRs), it is crucial to immobilize enzymes on supports while retaining their structure and activity. In a recent study, Liu<sup>60</sup> described a unique technique utilizing dicarboxylic acid halides as spacers for surface-initiated, covalent immobilization of enzymes onto cellulose microporous membranes. Sebacoyl chlorides, dicarboxylic acid halides, exhibit excellent reactivity with hydroxyl groups and amino groups, which are easily introduced onto membrane surfaces (Fig. 11). Trypsin and lipase were immobilized on the membrane surface while maintaining their molecular structure and activity substan-

Table 1 Influence of lignocellulosic (or cellulose)-based support materials on the stability and activity of proteins

| Biomass                            | Biomass-derived material  | Used enzyme           | Immobilization<br>method | Result/performance study   | Ref. |
|------------------------------------|---|-----------------------|--------------------------|--|------|
| Cellulose                          | Spin coating of protein NFC conjugates  | Alkaline phosphatase  | Conjugation              | No loss in activity at 21 °C and >20% remaining activity after a   | 17   |
| Lignocellulosic                    | Tendril-like carbon helices   | Cytochrome c          |                          | week of incubation at 37 °C<br>Enhanced structural stability<br>and >150% higher activity than   | 50   |
| Cellulose                          | СНМ   | Lipase                | Physical<br>adsorption   | native Cyt c 1.4 times higher specific activity, 41 folds half-life (at 45 °C), enhanced thermal and pH stabilities  | 51   |
| Lignocellulosic                    | Cellulose/lignin hydrogel beads   | Lipase                | Physical<br>adsorption   | 2.6, 1.2 folds higher relative and specific activity than lipase-cellulose bead system. >3 times higher thermal stability and against lower pH (i.e. 3)                          | 52   |
| Wood mimetic                       | Cellulose-biopolymer composite hydrogel beads   | Lipase                | Entrapment               | Improved reusability with >80% activity retention  | 53   |
| Cellulose                          | Cellulose based membranes<br>entrapping enzymes over the<br>electrode   | GOx                   | Entrapment               | The enzyme electrode was stable at for least 6 months. Protected the leakage of enzyme. Glucose detection up to $10 \mu M$ with response time of $\sim 10 s$                     | 55   |
| Cotton knit fabrics<br>(cellulose) | Cotton fabrics oxidized by periodate  | Cellulase             |                          | (1) Enhanced binding capacity of cellulase (>10.3 mg g <sup>-1</sup> ). (2) Enhanced cotton hydrolysis and cellulase potential for adsorption                                    | 56   |
| Cotton yarn<br>(cellulose)         | Cotton yarn oxidised with sodium periodate  | Trypsin               | Covalent binding         | Retained the initial activity after<br>60 days of storage in<br>physiological solution   | 57   |
| Cellulose                          | Cellulose-CBD   | β-Galactosidase       |                          | Reusability for milk sugar<br>hydrolysis till 40 cycles with<br>64% activity retention   | 59   |
| Paper wastes (PW)                  | $Fe_3O_4$ and chitosan functionalised $\alpha CFs$  | Laccase               | Covalent bonding         | Great loading capacity of >73 mg g <sup>-1</sup> ; ~92% activity recovery and excellent reusability capability with ~74% activity retention at the end of 11 <sup>th</sup> cycle | 61   |
| Waste newspaper                    | Dialdehyde-modified cellulose nanocrystals (DMC)  | Laccase               | Covalent bonding         | 64.94% yield, excellent stability, and reusability   | 62   |
| Cellulose                          | Magnetic dialdehyde cellulose<br>nanoparticles (MDC)  | Rhizopus lipase       | Cross linking            | Enhanced long term stability<br>with recovery rate >50% after 30<br>days   | 63   |
| Sugarcane bagasse                  | CNC   | Lipase                | Covalent bonding         | With optimum pH shifted to<br>8.25, found best suited to<br>catalyse a triglyceride lipolysis<br>reaction from palm oil  | 64   |
| Nanocellulose from<br>almond shell | Nanocellulose (NC) extracted using <i>p</i> -toluenesulfonic acid (PTSA) and sulfuric acid (ASS) with sugar-based natural deep eutectic solvent (NADES1a) as a biocatalyst system | CRL                   |                          | Significant improvement in half-life <i>i.e.</i> , 14 days greater than native enzyme  | 65   |
| Cellulose                          | Cellulose beads   | Glucose oxidase (GOx) |                          | Effective antimicrobial action against <i>Pseudomonas aeruginosa</i> , <i>Escherichia coli</i> and two of the methicillin-resistant <i>Staphylococcus</i> species                | 66   |
| Cellulose                          | Nonwoven cotton fabric  | Lysozyme              | Cross linking            | Enhanced antimicrobial activity and storage stability  | 67   |
| Cellulose                          | CMC-silver nanoparticle (AgNp)-silica hybrids   | Amylase               | Adsorption               | Activity increased and<br>maintained the activity at room<br>temperature (40 °C) for 15 days   | 68   |
| Cellulose                          | Cellulose powder, cotton buds,<br>disc make-up remover pads,<br>cotton and linen tissues  | Lipase                | Covalent bonding         | Increased effectiveness in<br>removing aged linseed oil layers<br>in 45 min at pH 6 and 40 °C  | 69   |

Table 1 (Contd.)

| Biomass   | Biomass-derived material   | Used enzyme  | Immobilization method                  | Result/performance study   | Ref. |
|---|--|--|--|--|------|
| Cellulose from coffee filter  | Cellulose nanofibers (CNFs)  | α-Chymotrypsin (α-CT)  | Enzyme<br>precipitate<br>coating (EPC) | Magnetic separation technique proved to be the best with >38 times higher activity than other methods. For long term, precipitate coating method proved to be superior with >70% activity retention even after 1 month of incubation | 70   |
| Cellulose   | TEMPO-oxidized cellulose fibres<br>used to build cellulose-based<br>microspheres | Phospholipase  | Covalent bonding                       | Showed a greater thermal<br>stability and resistance to pH, as<br>well as easy recovery and<br>reusability   | 71   |
| Cellulose   | Dialdehyde cellulose   | α-Amylase  | Covalent binding                       | Lys142 has been found to be involved in α-amylase immobilization to dialdehyde cellulose   | 72   |
| Cellulose   | Magnetic dialdehyde cellulose (MDAC)   | Bacterial laccase  | Cross linking                          | (1) With >210 mg g <sup>-1</sup> loading capacity and >34 emu g <sup>-1</sup> of magnetization, the material offers 10 cycles of reusability with >70%. (2) Efficient against crystal violet decolouration                           | 73   |
| Cellulose   | Microcrystalline cellulose (MCC) magnetic support                                | β-Galactosidase  |  | >90% immobilization efficiency,<br>1.2 folds higher substrate<br>affinity, 7 times higher thermal<br>stability and recyclability up to<br>15 cycles  | 74   |
| Nanocellulose (NC)<br>from an agro-waste<br>of quinoa husks<br>(QS) | Nanocellulose nano-carrier   | Laccase enzyme<br>(PersiLac1)  | Adsorption                             | Having ~98% and ~60% dye<br>degradation capacity pertaining<br>to Malachite green and Congo<br>red respectively, biocatalyst can<br>be reused with >83% initial<br>activity after 18 h   | 75   |
| Cellulose from Balsa<br>wood  | Balsa wood-derived cellulose<br>scaffold crosslinked with<br>glutaraldehyde      | Horseradish peroxidase<br>(HRP), glucose oxidase<br>(GOD), and catalase<br>(CAT) | Adsorption                             | >90% phenol degradation rate<br>and 95% sodium gluconate<br>yield were achieved in the<br>reactor  | 76   |

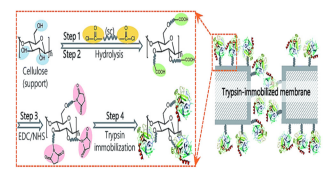


Fig. 11 Proposed pathways for trypsin immobilization on regenerated cellulose membranes. Reproduced from ref. 60 with permission from RSC, copyright 2017.

tially. The corresponding membrane demonstrated outstanding specific activity with strong activity retention (>25%) in a soaking mode at the optimal surface density. Moreover, the immobilized enzyme exhibited outstanding enhancement in reusability, thermal resistance, and continuous-operation capability.

Gajanan *et al.*<sup>61</sup> used paper wastes (PWs) to extract multifunctional  $\alpha$ -cellulose fibers ( $\alpha$ CFs), which were further tuned with magnetic Fe<sub>3</sub>O<sub>4</sub> and chitosan. This system was successfully employed for immobilization of laccase (Lac) with loading capacity of >70 mg g<sup>-1</sup> and >90% of activity recovery. Due to excellent pH, storage and temperature stabilities, authors claimed that the catalytic system could be applied for the degradation of DR28 (xenobiotic benzidine-based azo dye).<sup>61</sup> Overall, the unique microstructure of cellulose helps in improving the long-term stability and prevents back-diffusion of enzymes, which makes it an appropriate immobilization matrix for enzymes (Table 1).

### 3. Lignin-based supports (carriers) for enzyme immobilization

Lignin is a valuable and less explored natural resource. It is an intriguing class of material that has become a popular alternative in the search for more affordable and environmentally acceptable materials for enzyme immobilization. It possesses high strength and stability at extremes of pH, temperature, or

Table 2 Influence of lignin-based support materials on the stability and activity of proteins

| Biomass                           | Biomass-derived material  | Used enzyme                            | Immobilization method        | Result/performance study  | Ref. |
|-----------------------------------|---|--|------------------------------|---|------|
| Dendrocalamus<br>Latiforu         | BSS lignin as a support<br>material   | α-Amylase                              | Adsorption                   | With 19 mg g <sup>-1</sup> loading capacity, immobilized enzyme showed >2-folds higher activity with >53% activity retention after 14 recycles  | 86   |
| Bio-waste of kraft<br>pulp lignin | Amino-modified<br>microspheres (A-LMS)  | β-Galactosidase                        | Electrostatic interaction    | 1.5 times higher rate of galacto-<br>oligosaccharide production and efficient in<br>degradation of pesticide – lindane  | 87   |
| Green coconut<br>fiber (GCF)      | Lignin/Fe <sub>3</sub> O <sub>4</sub> nanoparticles synthesized by organosolv pre-treatment | β-Glucosidase                          | Covalent bonding             | (1) Improved digestibility performance (>21 g L <sup>-1</sup> of cellulose and >6 g L <sup>-1</sup> of green coconut fibre) and stability. (2) Good synergism with cellulases during enzymatic hydrolysis. (3) High adsorption capacity of nanoparticles towards Methylene blue (203.66 mg g <sup>-1</sup> ), Cibacron blue (112.36 mg g <sup>-1</sup> ) and Remazol red (96.46 mg g <sup>-1</sup> ) dyes | 88   |
| Kraft lignin                      | Novel hybrid support<br>made of titanium and<br>lignin                                      | α-Amylase from<br>Aspergillus oryzae   | Covalent bonding             | Improved pH range of 3–7 and best suited for low temperature catalytic applications (5–10 °C)   | 89   |
| Lignin                            | Lignin nanocapsules   | Tyrosinase and<br>laccase              |                              | Used for the synthesis of bioinks with<br>different shades. The nanocarriers found to<br>prevent alkaline and UV degradation of<br>synthesized melanin  | 90   |
| Organosolv lignin<br>(OL)         | LNPs  | Enzymatic cascade of lipase-tyrosinase | Layer-by-layer<br>method     | Successfully synthesised high yield lipophilic hydroxytyrosol esters  | 91   |
| Kraft lignin                      | Chitosan-coated LNPs  | Glucose oxidase<br>(GOx)               | Adsorption                   | Enhanced the enzyme activity up to 70° C<br>and display of self-scavenger property<br>towards H <sub>2</sub> O <sub>2</sub>   | 92   |
| Lignin                            | Lignin-based spherical particle   | Lipase                                 | Electrostatic<br>interaction | Retained 75% and 81% relative activity at higher temperature (60 °C) and after 10 reuses respectively   | 94   |

pressures (Table 2). 86-94 Weihua et al. 86 demonstrated the use of lignin particles from bamboo shoot shells (BSSs) as novel carriers for immobilization of  $\alpha$ -amylase from Bacillus subtilis with a maximum protein load of 19.0 mg g<sup>-1</sup> in 20 min. 86 The immobilized enzyme was superior compared with the free enzyme in terms of overall catalytic efficiency, storage stability, and recovery. Bebić and co-workers87 revealed amino-modified microspheres of kraft-lignin as potential supports for the immobilization of β-galactosidase and Lac enzymes. The former was used for the selective synthesis of galacto-oligosaccharides, whereas the latter was employed for the degradation of a pesticide (lindane). Lac immobilized onto modified kraftlignin showed 3-fold greater efficiency in degrading the pesticide than amino-modified silica nanoparticles.87 Furthermore, multifunctional lignin/Fe<sub>3</sub>O<sub>4</sub> nanoparticles were synthesized using green coconut fibres, which were used to remove colours from the effluents of the textile industry and to achieve a high stability for  $\beta$ -glucosidase.<sup>88</sup> Immobilized  $\beta$ -glucosidase revealed significant adsorption capacity for the elimination of methylene blue (203.66 mg g<sup>-1</sup>), remazol red (96.46 mg g<sup>-1</sup>), and cibacron blue (112.36 mg g<sup>-1</sup>). It also showed excellent digestion performance towards crystalline cellulose compared with that of the native enzyme.88 Therefore, the proposed approach proved successful in obtaining lignins isolated from lignocellulosic residues.88 Łukasz et al.89 designed a hybrid TiO<sub>2</sub>/lignin support for the covalent immobilization of Aspergillus oryzae α-amylase. The immobilized enzyme dis-

played improvement in chemical and thermal stabilities with 10 cycles of reusability, and after 1 month of storage its initial activity was retained at >80%. Considering sustainable and green synthetic protocols, Capecchi and colleagues<sup>90</sup> described a method of preparing sustainable bio-ink ingredient from poly(diallyldimethylammonium chloride) (PDDA) or chitosanfunctionalized organosolv lignin nanocapsules. Pigments were catalysed by Lac and tyrosinase immobilized over functionalized lignin nanocapsules *via* a layer-by-layer (LBL) immobilization strategy.

The lignin nanocapsules were employed as multifunctional devices acting as scaffolds, activators for enzymes to catalyse pigment synthesis and, finally, being the vehicle or binder for pigments. L-Tyrosine, epicatechin, 3,4-dihydroxyphenyl-Lalanine (L-DOPA), and several other derivatives of phenol were used as reactants giving out melanin pigment with high alkaline and UV tolerance due to interaction with nanocapsules. Similarly, Tomaino et al.91 synthesized one-pot esterification of lipophilic hydroxytyrosol esters by a lignin nanoparticles (LNPs)-supported enzymatic cascade. Tyrosinase and lipase M were immobilized on LNPs by an LBL method using chitosan as a positive-charged polyelectrolyte and concanavalin A acting as molecular spacer to regulate the optimal separation between the active sites of the two enzymes. Interestingly, compared with the native counterparts, both immobilized enzymes on LNPs showed greater kinetic parameters (maximum reaction speed  $(V_{\text{max}})$  and Michaelis constant  $(K_{\text{m}})$ ). By further

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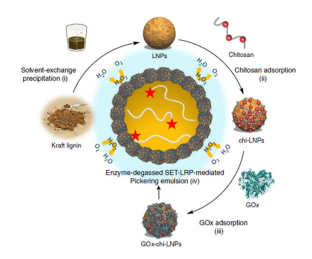


Fig. 12 Preparation of GOx-chi-LNPs and application in enzymedegassed SET-LRP-mediated Pickering emulsion (Schematic). (i) Preparation of LNPs via solvent-exchange precipitation. (ii) Adsorption of chitosan on LNPs to yield chi-LNPs. (iii) Adsorption of GOx on chi-LNPs to yield Gox-chi-LNPs. (iv) Application of GOx-chi-LNPs as biocatalytic degassing stabilizers for SET-LRP in Pickering emulsion media. Reproduced from ref. 92 with permission from Nature Portfolio, copyright 2020.

extending the LBL strategy, Moreno and colleagues<sup>92</sup> prepared a multifunctional particulate emulsifier where GOx was immobilized over chitosan-coated LNPs (Fig. 12).

This approach proved to be efficient in stabilizing the Pickering emulsions and in situ enzymatic degassing of single electron transfer-living radical polymerization (SET-LRP). Because of the auto-degradation of H2O2, GOx is protected against the oxidative stress caused in situ, leading it to withstand temperatures higher than the optimum. This rational design of the process is a breakthrough in avoiding the use of additives for SET-LRP and in opening new avenues for the green synthesis of block co-polymers having complex architectures. For the first time, applications of functionalized lignin as a matrix for immobilizing a wide range of enzymes (transaminase, carboxylase, dehydrogenase) using various binding chemistries were carried out by Benítez and co-workers.93 As a proof of concept, ω-transaminase was incorporated into a packed bead reactor and reversibly immobilized on polyethyleneimine-lignin, and its endurance was investigated in continuous-flow deamination reactions maintaining the same conversion for 100 cycles. All of these results demonstrated that fully closed-loop sustainable flow-biocatalytic systems based on bio-waste and natural lignin material to be effective substitutes for typical immobilization supports.

### Silk fibroin-based supports (carriers) for enzyme immobilization

Silk fibroin (SF) is a well-known and widely used natural protein procured from silkworm cocoons. Large numbers of

Fig. 13 Chemical structure and illustration of the possible hydrogen bonding (--) sites in silk fibroin.

serine, glycine and alanine residues are found in this macromolecular protein (Fig. 13), <sup>24,95,96</sup> along with readily reacting chemical groups (e.g., imidazole, tyrosyl/phenol and sulfhydryl).97 Due of its low cost, abundance, nontoxicity, and biodegradability, SF makes an appealing material for enzyme entrapment. Its peculiar amino-acid sequence and structure result in the formation of discrete nanoscale pockets and ensure that a sufficient number of water molecules is retained for protein interaction and stabilization, which make it a promising substrate for enzyme immobilization. 16 The distinctive structure of SF protein ( $\alpha$ -helices,  $\beta$ -sheets and random coils) synchronize to create a propitious space for stabilizing entrapped enzymes. SF can be transformed into fibres, powdery nanoparticles or microspheres, membranes, gels, films, hydrogels, or scaffolds under mild, ambient, or aqueous conditions. 90 Numerous studies have shown that SF can serve as a matrix for immobilization, and thereby to impart stability to a variety of enzymes.87-101 Recently, various works have been published addressing the use of silk biomaterials as excellent supports for the covalent and noncovalent immobilization of enzymes with high retention of catalytic activity and stability against various stresses (Table 3). 100-114 Silk nanofiber (SNF) is an excellent candidate for enzyme immobilization due to superior stiffness, larger specific surface area, higher interconnectivity and porosity with high capacity for loading enzymes. In this regard, Lee et al. 91 prepared SF nanofibers by electrospinning and applied it for immobilization of α-CT.

The activity of immobilized α-CT on SF nanofibers was shown to be eight-times more active than that of bulk material, and the activity was indirectly proportional to fibre diameter. 99 Many reaction sites (e.g., -NH2, -COOH, imidazole and phenol) are provided by the diverse amino-acid constituents of silk fiber. Lipase was immobilized onto SF using a chemical method. 100 This lipase was then used for the hydrolysis of Helianthus annuus (sunflower) oil for producing fatty acids. 100 Chen and co-workers 101 designed an inexpensive woven fabric support made from SF for the immobilization of Canadida sp. lipase. The stability and activity of immobilized lipases on two types of silk fabrics were examined, and it was shown that the lipase immobilized over the hydrophobic fabric had 2-fold higher esterification and hydrolysis activity than the lipase immobilized on native silk fibres. In comparison with free lipase, lipase after immobilization demonstrated stability over a larger pH range and a change in its optimum temperature. The method described in this work indicates that woven SF might be an ideal material for use as an immobilization

 Table 3
 Influence of silk-based support materials on the stability and activity of proteins

| Biomass                                  | Biomass-derived support material  | Used enzyme                          | Immobilization<br>method | Result/performance study  | Ref |
|--|---|--------------------------------------|--------------------------|---|-----|
| Bombyx mori<br>silkworm                  | Silk fibroin film   | GOx                                  | Entrapment               | Activity maintained more than 10 months of preservation up to   | 23  |
| cocoons<br>Bombyx mori<br>silkworm       | Silk fibroin film   | HRP                                  | Entrapment               | 37 °C of temperature<br>Even 2 months later, 90% of activity<br>retained at room temperature  | 23  |
| cocoons<br>Silkworm<br>cocoons           | Silk Fibroin nanofibers   | α-СΤ                                 |                          | High loading capacity of 5.6% with 8 times higher activity than the one immobilized on silk fiber and >90% activity retention after 24 h incubation       | 99  |
| Woven Muga<br>Silk                       | Glutaraldehyde – crosslinked silk<br>fibers   | Lipase                               | Cross linking            | 12% lower activity than native enzyme but significant conversion rates in the emulsion phase with 3 cycles of recycling                                   | 100 |
| Bombyx mori<br>silkworm<br>cocoons       | Hydrophobic silk fiber treated with polydimethylsiloxane (PDMS)   | Lipase                               | Adsorption               | 2-Fold increase in the activity and improved thermal stability  | 101 |
| Bombyx mori<br>silk                      | Silk fibroin crosslinked with tyrosinase using glutaraldehyde   | Tyrosinase                           | Cross linking            | (1) A higher degree of stability. (2)<br>80% of initial activity retained even<br>after multiple times reuses   | 102 |
| Woven Muga silk                          | Silk mat using <i>N</i> -ethyl- <i>N</i> '-(3-dimethylaminopropyl carbodiimide)<br>EDC-NHS ( <i>N</i> -hydroxysuccinimide)<br>linkage | ChOx                                 | Covalent<br>bonding      | Greater catalytic activity and<br>structural stability sustaining for<br>13 months and 25 individual assays   | 103 |
| Bombyx mori<br>silk                      | Silk fibroin-based hydrogel   | CA                                   |                          | (1) ~100% immobilization efficiency.<br>(2) Stable against pH (pH-3)<br>denaturation  | 104 |
| <i>Bombyx mori</i><br>cocoon             | Silk film by entrapment   | β-Glucosidase                        | Entrapment               | Enhanced stability against various stresses such as heat, electrodialysis, and protease   | 105 |
| Bombyx mori<br>cocoons                   | Silk film entrapment  | ОРН                                  | Entrapment               | (1) Excellent long term stability with >60% activity even after 17 weeks of incubation. (2) Higher stability against UV radiation and extreme temperature | 106 |
| Bombyx mori<br>silk cocoons              | Silk powder by entrapment   | Invertase                            | Entrapment               | Enhanced thermal stability  | 107 |
| Bombyx mori<br>silk cocoons              | Powder of Silk-ASNase bioconjugates<br>by glutaraldehyde  | ASNase                               | Bioconjugation           | Improved half life (63 h), 6 times<br>higher substrate affinity, resistance<br>to trypsin digestion and good<br>thermostability                           | 108 |
| Bombyx mori<br>silk cocoon and<br>lignin | Hydrogels composite made of silk fibroin and alkaline lignin (SF-AL)  | Laccase (Lac)                        | Cross-linking            | Enhanced working range of pH from 4.5 to 9  | 109 |
| Bombyx mori<br>silk cocoons              | SF nano-cocoons   | GOx and HRP                          | Encapsulation            | Improved the stability and efficiency<br>of the catalytic reaction, as well as<br>the temperature stability and storage<br>stability                      | 111 |
| Bombyx mori<br>silk cocoons              | Silk fibroin hydrogel   | Microalgae                           | Entrapment               | Good mechanical strength and<br>stability of material with 6 times<br>higher oxygen generation compared<br>to controlled algal culture                    | 112 |
| Bombyx mori<br>silk cocoons              | SF  | GOx                                  | Encapsulation            | Biocompatible and biodegradable<br>sensor with improved current<br>response of 0.2 μA mM <sup>-1</sup>  | 113 |
| Bombyx mori<br>silk cocoons              | SF  | Protease                             |                          | Good extraction efficiency  | 114 |
| Silk                                     | A film made of sericin, polyvinyl alcohol, and cassava starch   | Botryosphaeria<br>ribis EC-01 lipase | Adsorption               | >9.8 folds higher activity, better yield, and recyclability up to 7 cycles  | 115 |
| Bombyx mori<br>silkworm<br>cocoons       | Sensor-based on regenerated silk<br>fibroin film  | HRP                                  | Entrapment               | Improved detection limit of 100 nM with <40 s response time   | 116 |
| Bombyx mori<br>silkworm<br>cocoons       | Porous membranes of Bombyx mori silk fibroin  | GOx                                  | Entrapment               | Improved catalytic activity and<br>thermal stability with >20 times<br>increased permeability of NaCl and<br>glucose                                      | 117 |

Table 3 (Contd.)

| Biomass  | Biomass-derived support material                                      | Used enzyme                    | Immobilization method | Result/performance study   | Ref. |
|--|---|--------------------------------|-----------------------|--|------|
| Bombyx mori and<br>Philosamia<br>cynthia ricini silk | SF membrane   | GOx                            | Entrapment            | Improved shelf life with lower extent<br>of leakage (0.05%), and improved<br>thermal stability at 60 °C  | 118  |
| Woven silk   | Immobilization on silk Fibre using the diazo method                   | Ribonuclease                   | Adsorption            | 63% of activity retained for 7.2 months  | 119  |
| Bombyx mori<br>silk cocoons                          | Silk powder by entrapment   | Phenylalanine<br>ammonia-lyase | Entrapment            | Improved resistance against chymotrypsin and trypsin   | 120  |
| Bombyx mori<br>silk cocoons                          | Entrapment with silk blend<br>membrane and polyvinyl alcohol<br>(PVA) | GOx                            | Entrapment            | Improved catalytic activity  | 121  |
| Bombyx mori<br>silk cocoons                          | Graphene silk fiber nanocomposite decorated by platinum nanosphere    | GOx                            | Cross linking         | Stable conductivity of $\sim$ 57 S m <sup>-1</sup> with an improved sensitivity of 150 $\mu\text{A mM}^{-1}$ cm <sup>-2</sup> and detection limit of 1 $\mu\text{M}$ | 122  |

matrix in industrial preparations. 101 Similar to silk fibers, silk films offer handy and extremely efficient supports for the longstanding stability of enzymes that are confined. Lu and coworkers<sup>23</sup> investigated the stabilization of horseradish peroxidase (HRP), lipase, and GOx entrapped in self-assembled SF films with higher enzyme loading. Among the three enzymes studied, GOx retained significant activity over 10 months of shelf storage at a wider temperatures range until 37 °C. The extent of molecular interaction between enzyme molecules and SF chains, the susceptibility of the enzyme to undergo oxidation, and hydrophobic-hydrophilic interfaces have been postulated to influence the relative stabilizing outcomes among the three enzymes used in that study. 23 Acharya et al. 102 explored an alternative and efficient method for the biosynthesis of L-DOPA from tyrosine using tyrosinase immobilized onto a novel silk protein: fibroin. 102 Saxena and co-workers 103 further demonstrated silk mat produced by weaving silk fibres of Antheraea assamensis as a promising biocompatible matrix for cholesterol oxidase (ChOx). 103 The porous and fibrous structural morphology of the silk mat provided an ambient microenvironment for immobilizing ChOx with loading efficiency of >70%, which resulted in a good analytical display with higher sensitivity, stability, reproducibility and much greater selectivity for the substrate (cholesterol). On the other hand, protecting enzymes and proteins from undergoing inactivation induced by pH is a significantly difficult task. Taking up the challenge, Han and co-authors demonstrated that bombyx mori SF-based hydrogel with Ru(II)-mediated photo-chemical crosslinking served as an efficacious carrier for protecting an immobilized enzyme, carbonic anhydrase (CA), from pH denaturation. 104 The free enzyme lost its activity entirely at pH 3, whereas the immobilized CA maintained >20% of its initial activity. These results implied that CA was stabilized to a greater extent in SF-based hydrogels than the non-immobilized counterpart. Furthermore, the authors applied the same method for xylanase and lysozyme. They found that the immobilized enzymes achieved comparatively higher activity even at an unfavourable basic pH of 9.104 Silk membranes have also been found to impart stability to

enzymes against heat. Miyairi and colleagues 105 used a fibroin membrane as a support to immobilize β-glucosidase. Apart from heat tolerance, the immobilized biocatalyst could counter the effects of electrodialysis and protease treatments as well. 105 In addition, Dennis et al. demonstrated the value of SF entrapment for the preservation of OPH activity under a wide range of environmental circumstances, including high temperature, exposure to ultra violet light and to denaturing conditions imposed by organic reagents. 106 An enhancement in thermal stability similar to this work has also been reported by Yoshimizu<sup>107</sup> for SF powder over the enzyme invertase. In parallel, for treating acute lymphoblastic leukaemia, L-asparaginase (ASNase) is employed as a drug. However, to overcome its shortcomings in triggering an allergic response and a very short half-life, SF has been found to be suitable candidate for bioconjugation. 108 In this regard, to lower the immunogenic responses and the innate property of antigenicity of the enzyme, Zhang and colleagues 108 bioconjugated ASNase with polar functionalities of fibroin protein. The manipulated enzyme not only increased its stability against protease digestion (by trypsin), heat and storage, it also increased its half-life from 33 h to 63 h under laboratory conditions. 108 SF protein also offers a promising strategy to exploit the tremendous capabilities of Lac. 109 The adoption of gentle, aqueous processing techniques that were made possible by the specific chemistry, structure, and assembling of silk into nanodomains with low water content may be responsible for preservation of bioactivity during the manufacture of silk devices (Fig. 14).110

## 5. Chitin-based supports (carriers) for enzyme immobilization

Chitin is a biopolymer with potential applications in the biomedical sector. After cellulose, chitin is the second most abundant natural polysaccharide in the world. It is made up of  $\beta$ -(1–4) connected repeating units of *N*-acetyl-glucosamine (Fig. 15a) that can be found in the outer skeleton

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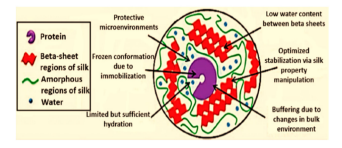


Fig. 14 Summary of the effects of stabilization of silk fibroin on immobilized compounds. Reproduced from ref. 110 with permission from Wiley-VCH, copyright 2012.

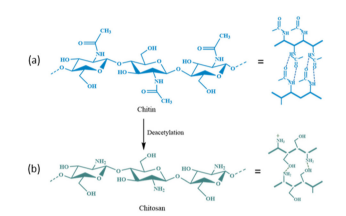


Fig. 15 Chemical structure and possible intermolecular hydrogen bonding (--) in (a) chitin and (b) chitosan.

of arthropods, insects, marine invertebrates, and cell walls of some fungi. 125-127 This biopolymer has many potential applications in medicine, food, cosmetic, agriculture, and immobilization processes due to its low toxicity and bioactivity, as well as high biocompatibility, biodegradability, and antibacterial properties. 128-136 The supports based on chitin and its derivatives provide enzymes with good stability and affordable bioprocessing, and they can be used in various applications, ranging from agriculture to drug delivery. The enzymes bound to the chitin matrix also have greater thermal activity than that of free enzymes. Table 4 130-150 displays examples of recent research that used chitin-based supports to enhance the stability and activity of enzymes.

Important factors for choosing an appropriate enzyme support include the degree of chitin deproteinization, the availability of amino groups, the presence of minerals, mesh size, surface structure, and its conformation. 131,132 The activity of invertase and amyloglucosidase has been improved by immobilizing them on chitin supports. 131 Xavier et al. 132 compared the characteristics of glucoamylase and α-amylase on two distinct types of support: ceramic and chitin. The initial activities of the glucoamylase complexes immobilized on chitin were five-times higher than those on ceramics, and the proportion of activity loss was two-times lower. 132 The physical properties of chitin are also dependent upon its source. For

example, due to the β-type crystalline structure, chitin from squid pens possesses superior physical qualities than chitin from crabs. 133 Chitin from squid pens was converted into a thin membrane that was immobilized with GOx. The membrane had the potential to entrap enzyme molecules as well as sufficient glucose and oxygen permeabilities to be beneficial for biosensors. The immobilized GOx on the chitin membrane was combined with an oxygen electrode to create a glucose sensor that had good stability and a wide linearity range (0.125-20 mM). 133 Immobilization methods can improved the properties of chitin further. To bind CRL, Gomes and coworkers 135 functionalized chitin with hexamethylenediamine followed by activation of glutaraldehyde. The activity profile as a function of pH, temperature, and thermal stability was determined for free and immobilized lipases. Authors revealed that the immobilized CRL had higher thermal stability than free CRL. 135 Leissy et al. 136 described a novel technique for immobilizing enzymes that relied on creation of polyelectrolyte complexes with supports coated in polymers with opposite charge. They used chitosan to chemically modify Saccharomyces cerevisiae invertase, which was immobilized further on a chitin support coated with sodium alginate. The protein-immobilized yield and activity was 85% and 97%, respectively. After immobilization, the optimal temperature and thermostability of invertase were increased by about 9-10 °C. The immobilized enzyme was four-times more thermally resistant than its native counterpart at 65 °C and stable against incubation in solutions of high ionic strength. 136 The authors suggested that the electrostatic immobilization technique they described might enhance the operational and functional features of the enzyme. 136 In another study, the same authors 137 described the immobilization of chitosan-invertase derivative on hyaluronic-acid-coated chitin. The immobilized enzyme was sixtimes more robust to thermal treatment at 65 °C than the native enzyme, and was stable against incubation in a solution of high ionic strength. Eighty percent of the initial activity of invertase was retained in the immobilized enzyme. The optimum temperature and thermal stability improved upon immobilization. The prepared biocatalyst showed exceptional temperature, storage, and operational-stability characteristics.137 The reusability and recovery of enzymes can be enhanced further by using magnetic-chitin (MCT) matrices, 138 wherein magnetic granules were synthesized using chitin as a protective and dispersive matrix. The easily synthesised MCT particles with an average size of ~1.5 µm were modified with dopamine to provide a useful matrix for immobilizing enzymes. Dopamine was self-polymerized and coated onto MCT to provide an enzyme-adhesive surface. An enzyme involved in starch hydrolysis, α-amylase, was immobilized on the polydopamine-coated MCT (DMCT). The effectiveness of enzyme immobilization on DMCT particles was boosted by glutaraldehyde treatment. 138 Over a longer range of pH and temperature, the surface-immobilized α-amylase showed comparatively greater activity than that of the free enzyme. Also, the glutaraldehyde-treated polydopamine functionalized MCT microparticles with immobilized α-amylase had excellent dura**Green Chemistry** 

 Table 4
 Influence of chitin-based support materials on the stability and activity of proteins

| Chitin<br>source                  | Chitin derived material  | Used enzyme                                      | Immobilization<br>method                  | Result/performance study  | Ref. |
|-----------------------------------|--|--|---|---|------|
| Chitin                            | Glutaraldehyde-activated chitin flakes   | Amylase  | Covalent bonding and crosslinking         | Superior operational stability with ~90% activity after 10 recycles and shelf life of >30 days with no loss in activity   | 130  |
| Chitin from<br>Krill              | Krill chitin   | Invertase and amyloglucosidase                   | Cross linking                             | Improved stability and activity   | 131  |
| Chitin from<br>squid pen          | β-Crystalline chitin membrane  | GOx  | Entrapment                                | The enzyme-immobilized membrane<br>is used for biosensors with wider<br>linearity range of 0.125–2 mM   | 133  |
| Chitin from crab shells           | Hexamethylenediamine<br>functionalized the chitin, and<br>activated it with glutaraldehyde           | CRL  | Covalent bonding                          | Improved half-life (>425 h), excellent thermal stability, great activity yield and protein retention  | 135  |
| Chitin from<br>lobster<br>shells  | Sodium alginate-coated chitin support  | Saccharomyces<br>cerevisiae invertase            | Electrostatic interactions                | 97% initial activity retained and good<br>thermal stability   | 136  |
| Chitin from<br>lobster<br>shells  | Hyaluronic-acid-coated chitin support  | Baker yeast invertase                            | Polyelectrostatic interactions            | Enhanced thermal stability, storage,<br>and operational stability. 69% of<br>enzyme activity retained at 37 °C<br>after 50 days of storage                        | 137  |
| Chitin                            | Poly dopamine coated magnetic chitin (DMCT) particles with glutaraldehyde                            | α-Amylase  | Cross linking                             | Immobilized α-amylase showed<br>higher durability, magnetic recovery<br>with 6 times reusability with >70%<br>activity retained                                   | 138  |
| Chitin                            | Immobilization of lysine decarboxylase (CadA) through fusion of a chitin-binding domain (ChBD)       | Lysine decarboxylase<br>(CadA)                   | Fusion of a chitin-<br>binding domain     | Better pH stability with >73% activity<br>at pH8 and high molar yield of<br>cadaverine (~97%)   | 139  |
| Chitin from<br>lobster<br>shells  | Chitosan-coated chitin support with glutaraldehyde   | Pectinase  | Adsorption                                | Greater stability and after nine<br>consecutive uses activity retained<br>100%  | 140  |
| shells of<br>shrimps<br>and crabs | CHNW   | Lysozyme   | Adsorption                                | (1) Enhanced antimicrobial activity against <i>Escherichia</i> , <i>Staphylococcus</i> and <i>Bacillus species</i> . (2) Improved enzymatic activity (>1.5 folds) | 141  |
| Chitin                            | Chitin and starch linked as a support  | Lipase and protease from sunflower seeds         | Covalent binding                          | Enhanced the efficiency of detergents stain removal   | 142  |
| Chitin<br>powder                  | AuNPs immobilized on CNFs  | GOx  | Not mentioned                             | Enhanced colorimetric glucose<br>detection with an LOD of 94.5 nM   | 143  |
| Chitin                            | Aerogel beads made of chitin/<br>graphene oxide (Ch/GO) composite<br>crosslinked with glutaraldehyde | CRL  | Adsorption<br>followed by<br>crosslinking | With immobilization capacity of >140 mg g <sup>-1</sup> , the biocatalyst can be reused for over 5 cycles with retention of 90% initial activity                  | 144  |
| Shrimp<br>shells<br>chitin        | Chitin   | Dextranase                                       | Adsorption and covalent binding           | A maximum of 88% immobilization<br>yield achieved with superior 22 folds<br>higher stability at 80 °C   | 145  |
| Chitin                            | Magnetic chitin nanofiber composite (MCNC)   | Chymotrypsin (CT)                                | Cross linking                             | Improved the loading capacity by 6.3 times after cross-linking, increased thermal stability and retained initial activity   | 146  |
| Chitin                            | Chitin powder  | α-Amylase from<br>Bacillus subtilis<br>ITBCCB148 | Adsorption                                | Immobilized $\alpha$ -amylase can be used up to five times and has an optimum temperature of 75 °C, increased the residual activity                               | 147  |
| Chitin                            | 3D fibrous chitinous scaffolds and silica nanopowder   | HRP  | Adsorption                                | Showed reusability of nano-SiO <sub>2</sub> (HRP)-chitin (HRP) scaffolds and high efficiency towards removal of 17-α-ethinylestradiol (EE2)                       | 148  |
| Chitin                            | Chitin powder  | Aspergillus Fumigatus<br>α-Amylase               | Adsorption                                | Improved thermal stability with >39% activity retained at 80 °C and 1.5 times higher activity at ambient conditions   | 149  |
| Chitin                            | Chitin-Bentonite Hybrid matrix   | α-Amylase from<br>Aspergillus fumigatus          | Physical adsorption                       | Enhanced stability with $\sim$ 4 folds higher $t_{1/2}$ compared to native enzyme   | 150  |

bility, magnetic recovery and reusability. 138 Zhou et al. 139 used chitin as the carrier support for efficient immobilization of CadA through fusion with a chitin-binding domain (ChBD) for the synthesis of cadaverine from L-lysine. 139 Furthermore, compared with wild CadA, the fusion protein (ChBD-CadA) had greater pH stability and maintained >73% activity at pH 8. 139 In a batch conversion, the immobilized ChBD-CadA (I-ChBD-CadA) converted L-lysine (200.0 g L<sup>-1</sup>) to cadaverine (135.6 g L<sup>-1</sup>), obtaining a 97% molar yield of the substrate (L-lysine). Furthermore, I-ChBD-CadA was reusable at high L-lysine concentrations and retained >57% of its initial activity after four cycles of usage without the addition of acid to maintain pH (Fig. 16). These results indicate that immobilizing CadA with a chitin-binding domain could be used for the industrial synthesis of cadaverine. 141 Ramirez and coauthors<sup>140</sup> developed a novel bioconjugate for immobilization of pectinase on a chitosan-coated chitin support. As a result of immobilization, heat and temperature resistance were improved and the enzyme remained stable with 100% of activity retained after nine consecutive uses, and ~70% of initial activity after 15 cycles of reuse. 140

Chitin-derived supports can also be used to enhance the antibacterial activity of various enzymes. Suisui et al. 141 investigated how chitin nanowhiskers (CHNWs) could enhance the antimicrobial and enzymatic activities of lysozyme adsorbed on CHNWs. Enzyme assays displayed that lysozyme-CHNW had 1.5-times higher enzymatic activity and greater antibacterial activity against Bacillus subtilis, Escherichia coli, and Staphylococcus aureus compared with those of the native lysozyme. These results revealed that lysozyme-CHNW could be used as a powerful antibacterial agent in food and medical fields. 141 Mehdi and co-workers developed a unique method for immobilizing protease and lipase on chitin-starch material as an enzyme biocatalyst matrix. Compared with native enzymes, immobilized enzymes had better pH, thermal, reusability, and storage stability. 142 Enzyme-immobilized chitincoated nanoparticles are effective biocatalysts due to their surface area and mechanical strength. 143,144 Yao et al. 143 presented a simple green method for the heterogeneous synthesis of stable, size-controllable gold nanoparticles (AuNPs)



Fig. 16 Immobilization of lysine decarboxylase for converting L-lysine to cadaverine *via* fusion of chitin-binding domains. Reproduced from ref. 139 with permission from Frontiers, copyright 2020.

immobilized on biocompatible chitin nanofibrils (CNFs). CNF-AuNPs showed peroxidase-mimic behaviour and catalysed the oxidation reaction of 3,3',5,5'-tetramethylbenzidine (TMB) by H<sub>2</sub>O<sub>2</sub>. When coupled with GOx, with a limit of 94.5 nM, CNF-AuNPs could detect glucose sensitively. This method had excellent specificity and efficiency, was affordable, and suffered negligible contamination, which showed its importance for the diagnosis of diabetes mellitus. 143 Chen et al. 144 fabricated chitin/graphene oxide (Ch/GO) composite aerogel beads for lipase immobilization using chitin as a great blending material. The synthesised beads had a three-dimensional porous structure, and GO was attached tightly to chitin with enhanced mechanical strength and surface area. Under ideal conditions, CRL was immobilized on Ch/GO aerogel beads. The immobilized CRL demonstrated excellent thermal stability compared with that of the native enzyme and, even after recycling five times, the immobilized lipase maintained initial activity of >90%. 144 Shahid et al. 145 improved the functionality of dextranase (isolated from a thermophilic bacteria Bacillus megaterium KIBGE-IB31) with the help of its immobilization on chitin by various protocols. Adsorption and covalent binding strategies were used to immobilize the isolated dextranase on chitin. Compared with the enzyme immobilized by an adsorption approach, dextranase immobilized by covalent cross-linking demonstrated maximum stability at high temperatures combined with improved recycling efficiency. Thus, chitin appears to be an inexpensive and convenient matrix for immobilizing a variety of enzymes to increase the stability and reusability of enzymes at an industrial scale.

### 6. Chitosan-based supports (carriers) for enzyme immobilization

Chitosan is a linear polysaccharide made up of N-acetyl-Dglucosamine (acetylated unit) and randomly arranged  $\beta$ -(1  $\rightarrow$ 4)-linked p-glucosamine (deacetylated unit) (Fig. 15b). 151,152 Chitosan can be used in medical applications due to its antibacterial, wound-healing, biodegradable, biocompatible, and non-toxic features. 153-155 The chitosan matrix can be researched as a support for enzyme immobilization. The storage, thermal, operational, and pH stabilities can be greatly improved for enzymes that are immobilized on the surface of chitosan (Table 5). 156-214 Chitosan-based materials can be produced in different geometrical configurations (e.g., microcapsules, membranes, beads/microspheres, coatings, fibers, gels, and sponges), which have shown good storage, thermal/ operational stability, and reusability properties. Owing to these flexible features, their capacity as carriers for a wide range of enzymes could be applied practically. Immobilizing xylanase in xanthan/chitosan matrices has revealed higher thermal stability and activity than those of native enzymes. 164 Çetinus et al. 167 demonstrated that catalase (CAT) immobilized into chitosan beads had enhanced resistance against denaturation due to pH or heat. Immobilized CAT had a Km value that was two-times greater than that of free CAT and its  $V_{\text{max}}$  decreased

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 Table 5
 Influence of chitosan-based support materials on the stability and activity of proteins

| Biomass  | Biomass derived material   | Used enzyme                                    | Immobilization<br>method                    | Result/performance study  | Ref. |
|--|--|--|---|---|------|
| Chitosan   | Magnetic gel microspheres made of chitosan, cellulose, and ${\rm Fe_3O_4}$                                     | GOx  | Cross linking                               | Higher conversion yield (>91%)<br>and great recyclability with >84%<br>activity retention even after 15<br>cycles                                   | 79   |
| Chitosan   | Mesoporous silica/titania with a chitosan coating  | β-D-Galactosidase                              | Covalent binding                            | High activity (1223 U g <sup>-1</sup> ), excellent efficiency (74%) and >90% activity retained after 15 cycles of reuses                            | 153  |
| Chitosan   | Chitosan beads activated with glutaraldehyde   | Lipase   | Not mentioned                               | 46.5% of initial activity was improved  | 155  |
| Chitosan   | Glutaraldehyde cross-linked chitosan<br>beads  | Pleurotus<br>nebrodensis WC<br>850 laccase     | Cross linking                               | Great decolourization<br>efficiencies for various reactive<br>and disperse dyes (>83–90%)<br>and >84% retained activity after<br>8 cycles of reuse  | 156  |
| Chitosan   | Manganese-ferrite nanoparticles coated with chitosan   | Laccase  | Covalent bonding                            | Best suited for pharmaceutical<br>waste degradation with 80%<br>removal rate for diclofenac   | 157  |
| Chitosan   | Genipin-activated chitosan support   | β-Galactosidase                                | Cross linking                               | Increased thermal and storage<br>stabilities with potential<br>application in food industries<br>harbouring lactose hydrolysis                      | 158  |
| Chitosan   | Chitosan matrix  | Bromelain, Papain<br>and Cysteine<br>proteases | Adsorption                                  | (1) Intact optimum pH and<br>temperatures. (2) ~5.8 times and<br>~7.6 enhanced stability for<br>Bromelain and Papain<br>respectively                | 159  |
| Chitosan   | Chitosan/organic rectorite composites  | Polyphenol<br>oxidase (PPO)                    | Physical adsorption<br>and covalent binding | (1) Higher enzyme activity<br>~8920–16370 U g <sup>-1</sup> . (2) 89–93%<br>phenol derivative removal within<br>2 h with 10 cycles of recyclability | 161  |
| Chitosan   | Chitosan nanoparticles   | L-ASNase                                       | Encapsulation                               | Significant decrease in the minimum inhibitory concentration (MIC) against microbes (5.26 mg mL <sup>-1</sup> )                                     | 162  |
| Chitosan   | Chitosan microbeads by microencapsulation  | α-Amylase and<br>invertase                     | Microencapsulation                          | Enhanced the stability and activity   | 163  |
| Chitosan   | Chitosan-xanthan hydrogels   | Endo-1,4-<br>β-xylanase                        | Noncovalent link                            | Enzymes show 60–70% higher activity than free enzymes   | 164  |
| Chitosan   | Copolymer of chitosan and poly-<br>glycidyl methacrylate (GMA)   | Urease   | Covalent bonding                            | Enhanced enzyme specific activity, temperature stability, pH stability, and storage stability   | 165  |
| Chitosan   | flakes and porous chitosan beads (PCB)   | Lipase   | Physical adsorption                         | Immobilized enzyme showed<br>better thermal stability at pH-6<br>and 40 °C  | 166  |
| Chitosan   | chitosan beads crosslinked with glutaraldehyde   | Bovine liver<br>catalase (CAT)                 | Cross linking                               | Improved enzyme thermal, operational, and storage stabilities   | 167  |
| Chitosan   | The chitosan and TMSO were coupled using the sol-gel method to create porous gels of chitosan-SiO <sub>2</sub> | GOx  | Entrapment                                  | Enhanced stability  | 168  |
| Chitosan from<br>dried<br>uttlebone<br>cartilage | Chitosan and glutaraldehyde-<br>crosslinked activated clay beads in<br>equal weights                           | α-Amylase,<br>glucoamylase and<br>β-amylase    | Cross linking                               | Superior operational stability<br>with 81% retention of activity<br>after 50 cycles of reuse  | 169  |
| Chitosan   | Magnetic chitosan microspheres cross-linked with glutaraldehyde  | Laccase  | Adsorption and cross-<br>linking            | Increased pH and temperature stability  | 170  |
| Chitosan   | Chitosan beads activated with glutaraldehyde   | PGA  | Covalent attachment                         | Good thermal and alkaline pH stabilities  | 171  |
| Chitosan<br>powder                               | Chitosan microspheres and sponges  | Cellulase                                      | Covalent method                             | Higher protein loading (~145 mg g <sup>-1</sup> ), better storage stability and reusability   | 172  |
| Chitosan   | Nanoparticles of chitosan made by the ionisation gelation process  | Neutral proteinase                             | Adsorption                                  | Enhanced operational, storage, and thermal stability  | 173  |
| Chitosan   | Nanoparticles of chitosan made by<br>the ionisation gelation process   | Neutral lipase                                 | Adsorption                                  | Increased the enzyme's efficiency by 13.17% compared to free lipase   | 174  |

| Biomass                        | Biomass derived material   | Used enzyme                                   | Immobilization<br>method       | Result/performance study  | Ref |
|--------------------------------|--|---|--------------------------------|---|-----|
| Chitosan                       | Chitosan beads (Ch-bead) were<br>attached to cibacron blue F3GA dye<br>(CB-Ch-bead)  | CAT   | Adsorption                     | Better thermal, operational and storage stabilities   | 175 |
| Chitosan                       | Chitosan beads   | Pepsin  | Cross linking                  | Increased thermal stability with shift in optimum temperature   | 176 |
| Chitosan                       | Polyvinyl alcohol (PVA) coated chitosan beads  | Cellulase                                     | Adsorption                     | from 40 to 50 °C<br>Immobilized cellulase showed<br>better yield, pH stability (at pH<br>7), good storage and operational<br>stability                                  | 177 |
| Chitosan                       | Chitosan nanofibrous membrane  | Lipase  | Entrapment                     | Improved storage stability with ~92% activity after 48 days of  | 178 |
| Antarctic krill<br>shells      | Glutaraldehyde-pretreated chitosan membranes   | Urease  | Entrapment                     | incubation<br>Improved thermal stability with<br>raise in optimum temperature<br>from 65 to 75 °C   | 179 |
| Cuttlebone<br>chitosan         | Chitosan-clay composite bead cross-<br>linked with glutaraldehyde  | β-Glucosidase                                 | Cross linking                  | Enhanced storage stability and activity in dried composite  | 180 |
| Powdered<br>chitosan           | Chitosan beads   | Chymotrypsin                                  | Covalent multipoint attachment | At 65 °C, the immobilised<br>enzyme's half-life increased from<br>0.57 hours at 55 °C to 7.8 hours  | 181 |
| Powdered<br>chitosan           | Chitosan and agarose with glutaraldehyde   | Candida Antarctica<br>lipase type B<br>(CALB) | Multipoint covalent attachment | Enhanced the thermal stability and activity   | 183 |
| Chitosan                       | Nanocapsules made of CKGM-CS   | L-ASNase                                      | Entrapment                     | Prevent the leaking. Better<br>stability and activity for broad<br>pH range   | 184 |
| Chitosan                       | Chitosan crosslinked with glutaraldehyde used for adsorption of $\text{Cu}(\pi)$   | CAT   | Cross linking                  | Though Cu-Ch-CAT had 3 folds lesser substrate affinity, but imparted greater thermal and storage stabilities compared to Ch-CAT   | 186 |
| Chitosan                       | Magnetic chitosan nanoparticles  | Laccase                                       | Entrapment                     | Increased storage stability and operation stability   | 187 |
| Pink shrimp                    | Modified chitosan beads  | HRP   | Covalent method                | Enhanced the thermal stability and operational stability  | 188 |
| Chitosan                       | Chitosan beads   | CA  | Physical adsorption            | Half-life increased by 24 h and<br>48 h compared to native enzyme<br>at –20 °C and R.T. respectively  | 189 |
| Chitosan from crab shells      | Chitosan-coated magnetite nanoparticles (Fe $_3$ O $_4$ -CS)   | Laccase                                       | Adsorption                     | Enhanced storage stability with >70% activity after 30 batches of   | 190 |
| Chitosan from                  | Chitosan macroparticles and  | β-Galactosidase                               | Adsorption                     | use 75–83% activity was retained  | 191 |
| shrimp shells<br>Chitosan      | nanoparticles<br>Glutaraldehyde crosslinked chitosan-<br>clay composite beads  | Tyrosinase                                    | Cross linking                  | even after 50 cycles of reuse<br>Optimum temperature, loading<br>effectiveness, and activity were<br>all increased  | 192 |
| Chitosan<br>powder             | N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimide (EDC) and N-hydroxysuccinimide (NHS) are utilised as coupling agents to covalently immobilise chitosancoated Fe <sub>3</sub> O <sub>4</sub> nanoparticles | Lipase  | Covalent method                | The stability remains same after 13 days of storage at 25 °C, the immobilised lipase exhibits enhanced operational stability, including wider temperature and pH ranges | 193 |
| Chitosan from<br>shrimp shells | Chitosan macrobeads by covalent crosslinking using glutaraldehyde  | Trypsin                                       | Covalent cross-linking         | Improved pH and temperature stability   | 194 |
| Chitosan                       | Chitosan-halloysite hybrid-<br>nanotubes cross-linking with<br>glutaraldehyde  | HRP   | Cross-linking                  | The immobilised HRP did not lose any activity after 35 days of storage  | 195 |
| Chitosan                       | Magnetic chitosan carriers   | Lipase and β<br>galactosidase                 | Entrapment                     | Demonstrated excellent long-<br>term stability without enzyme   | 196 |
| Chitosan                       | Glutaraldehyde-hardened alginate-<br>chitosan beads  | Inulinase                                     | Cross linking                  | leaking from the support At 50 °C gel beads showed excellent activity. In six days at room temperature, the immobilised enzyme retained 76% of its activity             | 197 |

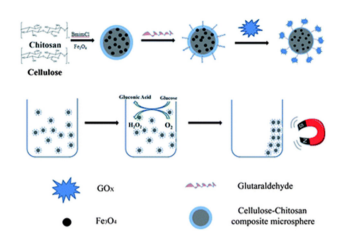
Table 5 (Contd.)

| Biomass   | Biomass derived material  | Used enzyme                           | Immobilization<br>method                              | Result/performance study   | Ref. |
|---|---|---------------------------------------|---|--|------|
| Chitosan  | ${ m Fe_3O_4}$ magnetic nanoparticles with chitosan coating and glutaraldehyde coupling agent   | Cellulase                             | Covalent method                                       | Greater operating stability<br>throughout a wider range of<br>temperatures and pHs, as well as<br>good activity and reusability<br>during magnetic separation<br>recovery                                    | 198  |
| Chitosan  | Hydrogel beads made of a bacterial cellulose (BC) and chitosan composite  | Lipase                                | Physical adsorption<br>and covalent cross-<br>linking | BC-chitosan hydrogel beads immobilized showed higher catalytic activity (1.9 times), and greater enzyme stability than microcrystalline cellulose MCC-chitosan hydrogel beads (22.7 times higher $t_{1/2}$ ) | 199  |
| Chitosan  | Dextran polyaldehyde is used as a<br>macromolecular cross-linking agent<br>to create chitosan magnetic<br>nanoparticles (CMNPs)                         | Pectinase                             | Cross-linking   | 2 folds improved thermal stability (55–75 °C) excellent stability and durability reusability   | 200  |
| Chitosan  | Chitosan-blended cellulose<br>monoacetate nanofibers  | Protease                              | Physical adsorption                                   | Higher operational stability, activity and reusability   | 201  |
| Chitosan  | Chitosan-montmorillonite<br>nanocomposite beads with<br>glutaraldehyde  | α-Amylase                             | Covalent bonding                                      | 95% activity retained after 40 days of incubation at 4 °C (~2.6 times higher than native enzyme)   | 202  |
| Chitosan  | Chitosan nanoparticles (CS-NPs)   | GOx                                   | Covalent attachment,<br>cross linking                 | Improved the stability and activity with ~26 folds higher activity (with precipitation method) compared to covalent attachment   | 203  |
| Shellfish<br>derived<br>chitosan (CS)<br>powder | Clay/chitosan biocomposite systems  | Protease                              | Covalent method                                       | Increased the immobilization yield   | 204  |
| Chitosan  | Magnetic chitosan nanoparticles (CS-Fe <sub>3</sub> O <sub>4</sub> ) modified with imidazole based ionic liquid (IL-CS-Fe <sub>3</sub> O <sub>4</sub> ) | Porcine pancreatic<br>lipase (PPL)    | Physical adsorption                                   | Having 1.93 folds higher specific<br>activity with 382% activity<br>recovery, immobilized PPL<br>showed 84% remaining activity<br>after 10 cycles of reuse   | 205  |
| Chitosan  | Chitosan-based nanoparticles  | Glucoamylase                          | Cross linking   | Greatly increased loading capacity and storage stability   | 206  |
| Chitosan  | Silica/chitosan composite   | Laccase                               | Covalent method                                       | Good residual activity after 7 months of storage   | 207  |
| Chitosan from crab shell                        | biocompatible glutaraldehyde (GA)<br>cross-linked chitosan beads as a<br>matrix   | Chitinase                             | Cross linking   | Immobilized enzyme was found<br>to be highly stable up to<br>1 month of storage and<br>exhibited wider range of pH<br>tolerance  | 208  |
| Cuttle-fish<br>waste                            | Highly swollen chitosan beads cross-<br>linked with glutaraldehyde  | Acid phosphatase<br>and β-glucosidase | Cross linking   | Enzymes were stable for long term  | 209  |
| Chitosan  | ${ m Fe_2O_3/chitosan}$ coated superparamagnetic nanoparticles was crosslinked with glutaraldehyde  | Lipase                                | Cross linking   | (1) Broader working temperature<br>range from 27–85 °C and pH<br>tolerance in between 5.5–9.5. (2)<br>Achieved 70.2% of microwave<br>assisted biodiesel recovery   | 210  |
| Chitosan  | Chitosan functionalized with Concanavalin A (ConA)  | α-Galactosidase                       | Adsorption, cross linking                             | Enhanced reusability and activity yield  | 211  |
| Chitosan  | Chitosan matrix   | Papain                                | Adsorption  | Enhanced stability against UV irradiation (up to 6040 J m <sup>-2</sup> )  | 212  |
| Chitosan  | Chitosan coated polylactic acid (PLA) based nanofiber (CCN) matrix  | α-Amylase                             |   | Higher substrate conversion ratio of 0.85–0.99 obtained with 40 min of residence time and lower rate of dilution   | 213  |
| Chitosan  | Chitosan gel activated with glutaraldehyde  | Cellulases                            | Cross linking   | Better hydrolysis performance<br>with Avicel and sugarcane<br>bagasse leading to glucose<br>concentrations of 10–13 g L <sup>-1</sup> in<br>the hydrolysate  | 214  |

from 32 000 µmol to 122 µmol (min mg protein)<sup>-1</sup>. However, immobilization enhanced the thermal, operational, and storage stabilities of CAT. 167 Yang et al. 168 immobilized GOx on a novel porous chitosan-SiO2 gel that was synthesized by coupling chitosan with tetramethoxysilane (TMOS) using a sol-gel method. With a profound immobilization yield of 97% and GOD activity of 1585 U g<sup>-1</sup>, even after 10 days and 15 days, the immobilized enzyme remained stable and functional, with activity of 86% and 56% at 30 °C, respectively, whereas most of the free enzyme became inactive after 5 days in a similar condition. 168 Mao and co-workers developed chitosan microspheres and sponges in smaller sizes to immobilize cellulase. 172 With rapid adsorption of cellulase over microspheres (<25 min), the immobilized cellulase displayed higher stability with respect to pH, higher K<sub>m</sub>, thermal stability, reuse, and storage stability than those of the native enzyme. Hence, this system could be employed in various industries demanding an enzyme with good performance at extreme physiochemical conditions. Dincer et al. 177 enhanced the stability of acid cellulase at neutral pH. Chitosan beads were coated with modified polyanionic PVA, and cellulase was immobilized on modified PVA-coated chitosan beads. Compared with the free enzyme, cellulase immobilized on chitosan beads demonstrated improved pH stability. The optimal pH of the enzyme moved from 4.0 to 7.0, signifying its ability for catalysis over a wide range of pH conditions, and also displayed excellent storage and operational stabilities. 169

To further explore the ability of a chitosan matrix to stabilize enzymes at different pH values, Tang *et al.* <sup>166</sup> prepared chitosan nanoparticles *via* ionization gelation, which yielded spherical nanoparticles of diameter 100 nm. Immobilized lipase showed improved activity, thermal, storage, and operational stabilities. The immobilized neutral lipase had a higher  $K_{\rm m}$  (0.37 × 10<sup>2</sup> g l<sup>-1</sup>) than that of free neutral lipase (1.01 × 10<sup>2</sup> g l<sup>-1</sup>). Moreover, immobilization improved the stability of neutral lipase in the acidic range. The stability features of the immobilized enzyme were enhanced noticeably with respect to temperature, reuse, and storage duration. <sup>166</sup>

To solve the problem of the separation and reusability of a biocatalyst, Liu and group<sup>79</sup> used the sol-gel transition method to develop magnetic Fe<sub>3</sub>O<sub>4</sub>-cellulose-chitosan hybrid gel microspheres by employing ILs as the solvent for the dissolution and regeneration of cellulose and chitosan. Glutaraldehyde was used to further immobilize GOx on hybrid gel microspheres (Fig. 17), which had a broader pH range, greater thermal stability, and enhanced storage stability than those of free GOx. The pH optimum of immobilized GOx was in the range 4-7, whereas that of free GOx exhibited maximal activity at pH 6. After 28 h of heating at 50 °C at pH 7, the immobilized enzyme exhibited 63% of its initial activity. However, under identical conditions, the free enzyme displayed only 22% of its initial activity.<sup>79</sup> Similar to the report stated above, Lac was immobilized on chitosan-coated magnetite (Fe<sub>3</sub>O<sub>4</sub>) nanoparticles. Kinetic analyses revealed that the immobilized Lac system had identical catalytic efficiencies. However, at optimal conditions, after 30 batches of use,



**Fig. 17** Preparation of magnetic Fe<sub>3</sub>O<sub>4</sub>-cellulose-chitosan microspheres after enzyme immobilization onto magnetic cellulose-chitosan (schematic). Reproduced from ref. 79 with permission from RSC, copyright 2012.

immobilized systems continued to exhibit >71% of their initial activity. 190 With regard to other closure studies, 188,191-196 lipase, β-galactosidase, cellulose, pectinase, and glucoamylase were immobilized onto chitosan-coated Fe<sub>3</sub>O<sub>4</sub> NPs. <sup>191-196</sup> These immobilized enzymes on magnetic chitosan carriers demonstrated prolonged stability with minimal leaching. They also conveyed better operational stability, including a wider thermal range, pH, reusability and storage stabilities, than those of the free enzyme. Hence, Fe<sub>3</sub>O<sub>4</sub>-chitosan nanoparticles could be used as efficient supports for lipase immobilization, particularly in applications involving the need for recycling. 188 Huang et al. 178 synthesised a chitosan nanofibrous membrane, and CRL was immobilized on the nanofibrous membrane using glutaraldehyde as a coupling reagent. Lipase loading on the nanofibrous membrane attained 63.6 mg  $g^{-1}$ , and immobilized lipase activity was 49.8%. After immobilization on the chitosan nanofibrous membrane, the stability towards temperature, pH, reuse, and storage of immobilized lipase was enhanced. With this good immobilization fraction of loading capacity and improved stability, the membrane of chitosan nanofibers is an excellent biocompatible and potential support for enzyme immobilization. <sup>178</sup> In the case pf porcine pepsin, immobilization onto chitosan beads shifted the optimum temperature by 10 °C, which was higher than that of the native version, revealing the resistance of the immobilized enzyme towards heat-induced denaturation along with a longer shelf-life. Observing these properties, Altun and colleagues<sup>176</sup> demonstrated the applicability of this system in industries based on the clotting and processing of milk. Adriano and co-workers<sup>181</sup> revealed a very noticeable improvement in the thermal stability of  $\alpha$ -CT after covalent bonding of the enzyme on a chitosan-based hybrid gel. The half-life of immobilized chymotrypsin was enhanced from 0.57 h at 55 °C to 7.8 h at 65 °C. 181 To improve the enzyme-holding capacity and to prevent leakage, Wang et al. 184 prepared carboxymethyl konjac glucomannan-chitosan (CKGM-CS) nanocapsules as a

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new biocompatible matrix system for the immobilization L-ASNase. The semi-permeability of the matrix retained L-ASNase in the matrix and inhibited leakage while allowing the substrate and product to pass through. The immobilized enzyme system demonstrated noticeably greater stability and activity in comparison with those of free L-ASNase. These investigations could offer a brand-new material for the immobilization of pH- and temperature-sensitive enzymes. 184 Pertaining to the challenge of reusability, Flores et al. 182 discovered a technique to activate chitosan beads using genipin, over which β-galactosidase from Aspergillus oryzae was immobilized. During batch processing, the immobilized enzyme maintained 100% of relative initial activity up to 40 cycles without much compromise in thermal stability, thereby showing its potential applicability. Similarly, a system of inulin over chitosan beads showed a recycling capacity up to 14 cycles after engineering by Singh et al. 185 Monier and co-workers<sup>188</sup> developed a method to immobilize HRP on modified chitosan beads by graft copolymerization of polyethylacrylate with potassium persulphate and Mohr's salt as redox initiators (Fig. 18). Even after immobilization, the optimum temperature remained at 45 °C with higher relative activity than that of the native enzyme. 188 Wanjari et al. 189 immobilized CA on chitosan beads and its activity was evaluated using the p-nitrophenol assay (Table 5). The storage stability was profiled for up to 20 days and the half-life of immobilized CA increased up to 48 h. The authors suggested that this system could be applied for carbon-capture and higher efficiency for converting CO<sub>2</sub> to CaCO<sub>3</sub>. 189 Zhai and colleagues 195 developed chitosan-halloysite hybrid-nanotubes (CTS-HNTs) through the assembly of chitosan onto hallovsite (natural nanotubular aluminosilicate). HRP was covalently immobilized over nanotubes through crosslinking (Fig. 19). Even 35 days later, the immobilized HRP maintained its full activity whereas the free enzyme could maintain only 27% of its initial activity. The authors claimed that the CTS-HNT-immobilized HRP system could be employed for wastewater rejuvenation, particularly for the destruction of phenolic chemicals, because of its strong catalytic activity. 195

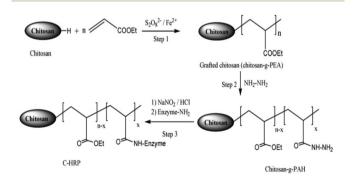


Fig. 18 HRP immobilization onto modified chitosan beads (schematic) Reproduced from ref. 188 with permission from Elsevier, copyright 2010.

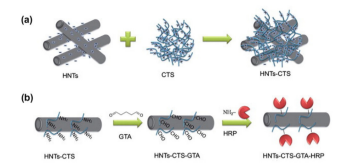


Fig. 19 (a) Stepwise preparation and (b) immobilization of HRP on a chitosan (CTS) matrix. Reproduced from ref. 195 with permission from Elsevier, copyright 2013.

### Alginate-based supports (carriers) for enzyme immobilization

Alginic acid is a naturally occurring polysaccharide and has a negative charge. It is extracted from algal cell walls, in particular brown algae or seaweeds. Alginate is a liner copolymer that consists of two residues: β-D-mannuronate (termed "M block") and α-L-guluronate (termed "G block"). Both residues are C5epimers (Fig. 20).<sup>215</sup> Depending on the source, the blocks M and G can be arranged in a consecutive manner (e.g., MMMM followed by GGGG or vice versa), alternating patterns (MGMGMGM), or random. 215 The properties of alginate (e.g., interaction with cationic metal species, solubility, and viscosity,) are directly attributed to the molecular weight of mannuronic acid/guluronic acid.215,216 Alginate is stable, biocompatible, biodegradable, nontoxic, has chelating ability, and relatively inexpensive. Thus, this natural polysaccharide and

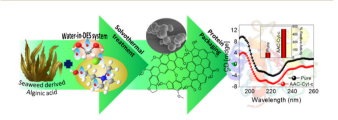
Fig. 20 Chemical structure of alginate. Reproduced from ref. 215 with permission from ACS, copyright 2005.

its derivatives have a multitude of industrial uses.<sup>216</sup> Alginate-based products have been used for drug delivery, regenerative engineering, environmental clean-up, wound dressing, biosensors, and transfection of genetic material.<sup>217,218</sup> An attractive approach to increase the stability of enzymatic processes and economic feasibility in terms of reusability can be achieved with the use of alginate-based supports for enzyme immobilization.<sup>219–226</sup> The mechanical characteristics and robustness of alginate-based supports can be enhanced further using polymer blending (much like cellulose and chitosan). These alginate forms have been demonstrated to increase thermal stability, enzyme activity, and reusability.<sup>227–236</sup>

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Palmieri et al.<sup>230</sup> developed a novel procedure for enzyme immobilization by entrapment in copper alginate gel, which was applied for immobilizing fungal phenol oxidase. The immobilized enzyme boosted the rate of oxidation of various aromatic substrates, was active over a large pH range, and had a low temperature for optimal activity. Thus, a system for detoxifying wastewater contaminated with phenolic derivatives from industrial or agricultural sources could be developed using an enzyme that has been immobilized in this way. 230 By first combining polysaccharides with Ca2+ and then crosslinking glutaraldehyde with the amine groups of the gelatine found in the initial mixture, Elçin and colleagues<sup>231</sup> created urease-containing xanthan-alginate spheres. Higher enzymatic activity was shown by encapsulated urease even with variations in pH or temperature. Even after 20-times of its use in optimal conditions, xanthan-alginate spheres continued to have 75% of the maximum urease activity.<sup>231</sup> More recently, a multifunctional alginate-derived carbon (AAC) material having variable oxygen functionalities was created by our research team<sup>232</sup> for the immobilization of Cyt c. Without affecting the structural stability if Cyt C, the enzyme activity was increased significantly in comparison with that of the native protein (Fig. 21).

For the entrapment of different enzymes, calcium alginate (Ca-ALG) use seems to be economical and biocompatible. Blandino  $et~al.^{233}$  systematically studied the activity of encapsulated GOx within Ca-ALG gel capsules. Comparison of the apparent kinetic characteristics of immobilized and free GOx revealed that the immobilized GOx had a higher  $K_{\rm m}$  and more pronounced differences in  $V_{\rm max}.^{233}$  Tanriseven and coauthors<sup>235</sup> immobilized Saccharomyces~cerevisiae invertase in alginate capsules, which achieved 87% relative activity along with enzyme stabilization at high pH and temperatures.<sup>235</sup>



**Fig. 21** Preparation of functionalised solvothermal carbon derived from alginate using "water-in-deep eutectic solvents" for enhancing enzyme activity (schematic). From ref. 232.

Alginate-based gels have also enhanced the thermal and storage stabilities of various proteins. 236 Vu and colleagues 236 investigated the biochemical characteristics of invertase entrapped within alginate gels. The  $K_{\rm m}$  of immobilized invertase (139.19 mM) was higher than that of free invertase (93.19 mM), but its  $V_{\rm max}$  was smaller. Nevertheless, the support induced greater thermal stability to invertase (as manifested by a longer half-life) and endowed significant longterm stability (up to 40 days). 236 To study the normalized effect of alginate matrices, Fadnavis et al. 237 employed lipase from porcine pancreas, Pseudomonas cepacia, and Candida rugosa. Hydrogels prepared by crosslinking a blend of sodium alginate (5%) and gelatin (3%) with glutaraldehyde were used to immobilize these lipases, which had excellent efficiency and greater stability and reclaimability for ~10 recycles without much loss of enzyme activity. Then, using a straightforward procedure of protein binding at pH 5 and release at pH 8.5, these functionalized beads were employed to purify crude porcine pancreatic lipase 7.4-times, showing their feasibility for biomedical applications.237 Thanks to advances in coreshell chemistry within nanoscience, Tagieddin and coworkers<sup>238</sup> designed alginate-chitosan core-shell microcapsules to create biocompatible carriers for enzyme immobilization. These shells offered permeability control over substrates and end products, whereas the protein was maintained in a liquid or solid core. To create a liquid or solid core for the microcapsule, sodium alginate was crosslinked with calcium or barium ions, respectively. o-Nitrophenyl-β-D-galactopyranoside (ONPG) was used to assess the catalytic activity of a model enzyme, β-galactosidase, which was immobilized in the alginate core. Due to the extra layer required for the influx of the substrate and outflow of the product,  $V_{\text{max}}$  for  $\text{Ca}^{2+}$  and  $\text{Ba}^{2+}$ alginate-chitosan core-shell microcapsules was much lower than that of the free enzyme. 238 Ultrathin alginate/protamine/ silica (APSi) composite membranes were constructed by Wang et al.<sup>239</sup> using a co-extrusion minifluidic and biosilicification methodology for Lac immobilization. The prepared hybrid capsules were highly monodisperse and dispensed Lac with significantly high thermal, storage, and pH stabilities. With ~100% immobilization yield, Lac displayed an activity of 61.8 mmol  $g^{-1}$  min<sup>-1</sup>, along with withholding 67% activity after 20 days and residual relative activity in APSi capsules remaining at 45% after 10 cycles.<sup>239</sup> Alginate-based hybrid composites have become an innovative class of materials, providing a larger range of uses for enzyme encapsulation encompassing biocatalysis, biomedicine, bioseparation, and biosensing. 240-245 To immobilize Lac, Lu et al. created alginate-chitosan microcapsules using emulsification-internal gelation.<sup>240</sup> In addition to stability, a higher loading efficiency and immobilized yields were obtained under ideal immobilization conditions (0.3% chitosan, 2% sodium alginate, 2%  $CaCl_2$ , and a 1:8 ratio by volume of enzyme:alginate). However, the recyclability was only three cycles, so finding a remedy for this problem is an issue.240 A multidimensional approach using biomass and nanomaterials to prevent leakage of yeast alcohol dehydrogenase (YADH) from Saccharomyces

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cerevisiae source was availed by Xu et al.  $^{241}$  By adding silica nanotubes (SiNTs) to the alginate (ALG) gel and then crosslinking it with calcium, they designed an alginate-silica nanotube (ALG-SiNT) composite, which was used to encapsulate YADH. Enzyme leakage from the carrier was reduced significantly ( $\sim$ 50%) compared with the alginate matrix composite only. These results showed that an ALG-SiNT carrier furnished stronger affinity with the enzyme compared with YADH immobilized over simple ALG support and led to longer storage, increased activity retention, and a stable operation.  $^{241}$ 

We applied a new and straightforward approach to prepare stable and versatile alginate-based white light-emitting hydrogels (WLEs) for protein packaging.242 Using the notion of complementary colors, a WLE composite was constructed by engineering the surface of an orange light-emitting ZnS quantum dot (QD) doped with Mn2+ using a blue light-emitting IL-choline-tosylate. The WLE QD-IL composite was combined with an alginate bio-polymer to fabricate the WLE hydrogel. After that, Cyt c was confined within the WLE hydrogel matrix, which led to higher relative activity of ~180% compared with that of the free enzyme (Fig. 22). The authors counteracted the harsh effects of chemical denaturants (e.g., urea) and extreme temperature over the stability of the metalloprotein Cyt c. Extending this concept further, Ma et al. 243 discovered an efficient carrier, double walled carbon nanotubedoped alginate gel (DWCNT-ALG) for immobilization of lactate dehydrogenase (LDH). Leakage of LDH from LDH-DWCNT-ALG biocomposite was decreased significantly, and LDH that had been immobilized displayed increased activity. The authors demonstrated the practical use of this type of carrier for lactate bioconversion in industry, as well as its usage for the immobilization of additional enzymes and microorganisms (including LDH and ADH).<sup>243</sup> Enzyme entrapment in ALG beads has been shown to be relatively inexpensive, safe, and straightforward technique for separation

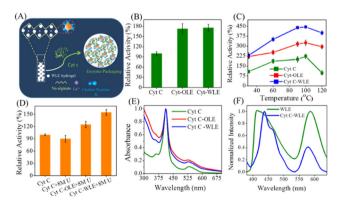


Fig. 22 Alginate-based hydrogels for protein packaging with enhanced activity and stability of Cyt c. (A) Schematic of the interaction of Cyt C with hydrogel composites. Percentage relative activity of Cyt C at room temperature (B), at various temperatures (C), and in presence of 8 M urea (D). (E) UV-vis spectra of Cyt C in presence of WLE and OLE hydrogels. (F) Fluorescence spectra of only WLE hydrogel and Cyt C + WLE hydrogels. Reproduced from ref. 242 with permission from Wiley-VCH, copyright 2020.

of the product and enzyme at the industrial scale.<sup>244</sup> Bhushan et al. 244 prepared ALG beads in an aqueous mixture containing sodium alginate, lipase from Arthrobacter sp. (ABL), and CaCl<sub>2</sub> to empower its reusability and all-round enzyme stability. In comparison with the free enzyme, the entrapped enzyme was much more stable throughout greater temperature domains, pH, and storage times. 244-248 Viet and colleagues immobilized cellulase in Ca-ALG beads by entrapment and its activity was assessed using CMC. With several cycles of use, the immobilized enzyme exhibited greater stability in response to changes in pH and temperature. The activity of immobilized cellulase remained 69.2% after five recycles and 20.3% after eight recycles. 246 Manganese peroxidase (MnP), generated from an indigenous variant of Ganoderma lucidum IBL-05, was immobilized on Ca-ALG beads by Bilal and co-workers<sup>247</sup> using a promising and environmentally friendly entrapment method. Ca-ALG-bound MnP was catalytically more vigorous and could be used eventually to decolorize dyes and detoxify industrial effluents more effectively.247 Nunes and coworkers<sup>248</sup> developed conjugated polyvinyl alcohol-alginate (PVA-ALG) beads with excellent thermochemical and mechanical stabilities at high temperatures (>80 °C). Naringinase, an enzyme derived from Penicillium decumbens, was immobilized in ALG (0.2-1.0%)-PVA (10%) beads featuring three sizes (1-3 mm). At pH 4.0 and 70 °C, immobilized naringinase bestowed 80% activity and displayed >90% retention of initial activity after 6 weeks of incubation in acetate buffer. The importance of this immobilization approach for the system under consideration was illustrated by these promising outcomes, which also implied that it could be used to entrap other biocatalysts.<sup>248</sup> Bonine et al.<sup>249</sup> reported immobilization of a novel lipase isolated from the seeds of Pachira aquatica (PAL) using Ca-ALG beads and PVA. Similar results were obtained with immobilization of PAL in ALG and ALG/PVA beads, with improvement in their stability against temperature than the free enzyme.<sup>249</sup> Ortega et al.<sup>250</sup> covalently immobilized neutrase on ALG-glutaraldehyde beads. Under optimized conditions (pH 6.2, 2% ALG, 6.2% glutaraldehyde, and interaction for 60 min), the immobilization yield of encompassed neutrase (61.84 U mL<sup>-1</sup>) was  $\sim$ 50% and the confined enzyme exhibited maximum activity at 10 °C higher than that of the native enzyme, which clearly showed the stability induced by the support. Immobilization caused the estimated activation energy to drop from 47.7 kJ mol<sup>-1</sup> to 22.0 kJ mol<sup>-1</sup>, which suggested the role of the support in favouring of the forward reaction at the thermodynamic level. 250 Eldin et al. 251 extended the covalent immobilization technique towards glucoamylase to form a novel affinity with p-benzoquinone-activated ALG beads. The covalently mounted enzyme sustained its activity after 30 successive runs and shelf space of 36 days. 251 To improve the stability and catalytic rate of GOx, Wang and coworkers<sup>252</sup> designed a protocol for undertaking emulsificationinternal gelation followed by GOx adsorption and chitosan deposition to encapsulate the enzyme in CaALG-chitosan composite microspheres (CACMs). Before grafting with chitosan, the Ca-ALG matrix was made highly porous with the release of

CO<sub>2</sub>, which helped to enhance the adsorption efficiency of the carrier. The concentration and molecular weight of chitosan, incubation duration, and pH seemed to have an impact on the GOx loading, encapsulation efficiency, and activity of CACM-GOx. Exhibiting the maximum enzymatic activity and encapsulation efficiency at an isoelectric point of 4, immobilized GOx retained >70% activity, which was tenfold higher than the activity retention by the free enzyme. 252 With some modification of this work, Zhao and colleagues<sup>253</sup> reported a unique and feasible approach for non-covalent GOx immobilization in chemically reduced graphene oxide (RGO)/ALG hybrid gel beads. The enzyme contained in the hybrid microbeads demonstrated high environmental tolerance and could maintain optimal activity over a wide range of conditions (45-60 °C and pH 4-6). To produce a continuous fixed-bed enzyme catalytic process, the microbeads could also be recycled simply through conventional filtering and reintroduced to a column.<sup>253</sup> Both of these works revealed the great potential to further explore CACM-GOx and ALG/RGO for developments in glucose bio-sensing. Abd El-Ghaffar et al. 254 used a grafting-encapsulation approach for  $\alpha$ -CT immobilization. First, chitosan grafted with polymethyl methacrylate (PMMA-g-CS) through free-radical polymerization was used as a support for α-CT. Ca-ALG beads were fabricated for sheathing PMMA-g-CS-CT to produce composite beads. For immobilized \( \alpha \) CT, greater retention of activity (~97.7%) was attained at pH 9 for 24 h. Immobilized α-CT retained 75% of its original activity after 60 days of storage at 25 °C and continued to work very well even after 25 reuses. Such results suggested that the recycling ability of ALG composite carriers could be used for continuous catalytic reactions in the industrial sector.<sup>254</sup> Muthuvelu and colleagues<sup>255</sup> designed a tri-enzyme biocatalyst by co-immobilizing Lac, cellulase, and β-glucosidase with ALG for evaluating bioethanol. The co-immobilized enzyme system had long shelf-life and thermal stability, offering an effective one-pot pre-treatment protocol for the production of bioethanol from lignocellulosic biomass, of which the maximum ethanol conversion was obtained with Ipomoea carnea (Fig. 23).

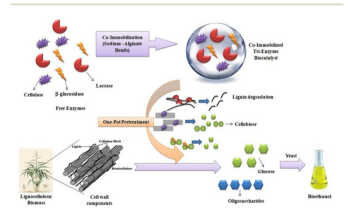


Fig. 23 Immobilization of co-immobilized laccase, cellulase, and  $\beta$ -glucosidase for ethanol production. Reproduced from ref. 255 with permission from Elsevier, copyright 2018.

nanofibers Electrospun comprising ALG-supporting immobilized lipase showed high enzyme loading capacity, catalytic activity, and stability. Doğaç et al. 256 produced PVA/ ALG and polyethylene oxide/alginate (PEO/ALG) nanofibres using electrospinning, and lipase was adsorbed and crosslinked further with glutaraldehyde. Free lipase lost all of its activity at high temperatures after 40-60 min, but lipaseimmobilized nanofibers sustained approximately 65-70% activity during the same timeframe. Furthermore, PVA/ALG and PEO/ALG composite nanofibers immobilized with lipase exhibited 50% of their original activity after 14 and seven cycles, respectively. 256 Zhao et al. 257 synthesized novel nanoflower/ALG microbeads and immobilized α-acetolactate decarboxylase (ALDC) using a facile method. First, ALDC was co-precipitated with Ca<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub> to form enzyme-inorganic composite nanoflowers (ALDC-Ca<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub>). Later, these nanoflowers were encapsulated in ALG gel beads (ALDC-Ca<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub> + ALG). <sup>256</sup> The microbeads demonstrated outstanding stability and environmental endurance in contrast to free ALDC and ALDC-Ca<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub> nanoflowers, displaying optimal activity over a wide range of temperature (45-70 °C) and pH (3.5-7), but retained >98% of activity in comparison with unbound ALDC. Furthermore, the immobilized enzyme was employed in a 300 L beer fermenter to stop diacetyl generation. This resolved the issue of off-flavor in beer and reduced the time it takes for beer to mature, thereby demonstrating enormous scope for application in breweries.<sup>257</sup> Xylanase has several uses in the food industry. Kumar et al. 258 purified xylanase from Bacillus licheniformis Alk-1, and entrapped it within crosslinked Ca-ALG beads, and undertook activation through glutaraldehyde. In comparison with the free form, immobilized xylanase demonstrated improved overall chemical characteristics, recycling, and reusability efficiency. Hence, xylanase-ALG beads with higher xylanolytic activity and stability could be prepared, thereby aiding the design of efficient bioreactors for various applications (e.g., formulations of poultry feeds). 258 Using citric acid as a nontoxic crosslinker, 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDC) and N-hydroxysuccinimide (NHS) as activators, Bedade and coauthors<sup>259</sup> mounted acrylamidase from *Cupriavidus* oxa laticus ICTDB921 on chitosan-coated Ca-ALG beads. After four cycles, the immobilized acrylamidase maintained 80% of its activity while exhibiting increased pH, temperature, and shelf stability.259 Pectic compounds cause turbidity in juices. The aesthetics and storage stability of such products can be enhanced by removing these substances with the help of pectinase.

Mohammadi and co-workers<sup>260</sup> immobilized *Aspergillus aculeatus*-originated pectinase on ALG-montmorillonite beads. At extremes of pH, immobilized pectinase showed higher efficacy than that of the free enzyme. After six successive cycles, immobilized pectinase exhibited >53% of its initial activity, demonstrating improved stability and renewability for pectin hydrolysis. Pineapple juice was clarified using this catalytic system, demonstrating the potential of this enzymecarrier model for use in fruit juices.<sup>260</sup> Table 6 summarizes the immobilization of other enzymes on the ALG matrix due to

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| Biomass              | Biomass derived material  | Used enzyme                          | Immobilization<br>method   | Result/performance study  | Ref |
|----------------------|---|--------------------------------------|----------------------------|---|-----|
| Alginate             | Polyacrylamide alginate cryogel<br>(PAG) functionalized with glycidyl<br>methacrylate (GMA)   | Laccase                              | Covalent<br>immobilization | >70% removal of phenolics from<br>olive mill wastewater, >55% dye<br>removal from textile wastewater and<br>93–99% decolouration of few other<br>dyes were achieved | 219 |
| Alginate             | Carrageenan-alginate beads  | CAT                                  | Covalent<br>immobilization | Enhanced pH tolerance towards alkaline media and 6 folds higher $V_{ m max}$ achieved   | 220 |
| Alginate             | sodium alginate (SA)-polyethylene glycol (PEG)  | Cellulase                            | Cross linking              | With only 22.68% immobilization<br>yield, 133% of yield (reducing sugar)<br>obtained  | 221 |
| Alginate             | Polypyrrole/silver nanocomposite coated with calcium alginate   | Polygalacturonase (PG)               |                            | Excellent immobilization efficiency (>84%) and long term stability (83% of initial activity after 2 months)   | 222 |
| Alginate             | Silver-alginate nanoparticle matrix   | Lipase                               | Conjugation                | Application in high-fat meat processing is demonstrated with loss of 40% initial weight after 2 weeks   | 223 |
| Alginate             | Dopamine-alginate beads   | Laccase from<br>Streptomyces cyaneus |                            | Amido Black 10B, Reactive Black 5,<br>Evans Blue, and Remazol Brilliant<br>Blue were all completely decoloured<br>with 100% efficiency                              | 224 |
| Alginate             | Alginate micron and submicron beads   | Alcalase                             | Covalent<br>immobilization | Improved performance and utilised<br>in 7 subsequent cycles of soy<br>protein hydrolysis with a slight<br>decrease in activity                                      | 225 |
| Alginate             | Sodium alginate (SA)-polyethylene<br>glycol (PEG)-Chitosan (CS)   | Cellulase                            | Entrapment                 | Increased stability and recyclability<br>with 22.68% activity higher than<br>native enzyme after 5 cycles   | 226 |
| Alginate             | Oxidized tyramine-alginates micro beads   | HRP                                  | Encapsulation              | With 20 mol% of tyramine-alginate, 96% of the phenol was eliminated from the solution. Increased reusability  | 227 |
| Alginate             | Carrageenan-alginate beads with chitosan-glutaraldehyde activation  | β-D-Galactosidase                    | Covalent<br>immobilization | Reusability was improved, and 79.74% of the activity from the first round maintained till the tenth reusability round   | 228 |
| Alginate             | Calcium alginate hydrogels with liposomes   | Bovine Carbonic<br>Anhydrase (BCA)   | Entrapment                 | Entrapment of BCA in alginate<br>hydrogels is stable and effectiveness<br>for using to catalytic reactions in<br>bioreactors  | 229 |
| Alginate             | Entrapment in copper-alginate beads   | Fungal phenol oxidase                | Entrapment                 | Enhanced thermal activity and<br>stability. Wider pH ranges allowed<br>the enzyme to remain active, and<br>storage at 4 °C considerably<br>enhanced                 | 230 |
| Alginate             | The gelatin component contained in the gel solution was cross-linked with glutaraldehyde (GA) and encapsulated in xanthan-alginate (XA) spheres | Urease                               | Encapsulation              | Increased temperature and pH<br>stability. Even after 20 times of use,<br>the xanthan-alginate spheres still<br>had 75% of their initial activity                   | 231 |
| Alginate             | Calcium alginate gel capsule<br>encapsulation   | GOx                                  | Encapsulation              | Studied gelation conditions affected<br>on capsule factors like capsule<br>thickness, enzyme leakage rate, and<br>encapsulation effectiveness                       | 233 |
| Alginic acid<br>(AA) | AAC   | Cyt c                                |                            | Enhanced enzymatic activity up to 5.5-fold and improved the thermal stability   | 232 |
| Alginate             | Calcium alginate gel capsule encapsulation  | GOx                                  | Encapsulation              | Studied gelation conditions affected<br>on capsule factors like capsule<br>thickness, enzyme leakage rate, and<br>encapsulation effectiveness                       | 233 |
| Alginate             | Encapsulation in microcapsules alginate coated with alternating   | Cyt c                                | Encapsulation              | Increased the encapsulation yield, activity as well as stability  | 234 |

invertase

Saccharomyces cerevisiae Entrapment

Ca-ALG gel capsules

alginate coated with alternating

multiple layers membrane of poly N-vinylamine and polyacrylic acid

Alginate

Relative activity was found 87% and 235

active for 36 days. More stable at high pH and temperatures

activity as well as stability

Table 6 (Contd.)

| Biomass   | Biomass derived material  | Used enzyme                           | Immobilization<br>method                          | Result/performance study   | Ref |
|---|---|---------------------------------------|---|--|-----|
| Alginate from Sargassum                               | Entrapped in alginate gel   | Invertase                             | Entrapment  | 3.5 times higher $t_{1/2}$ than native enzyme at 60 °C   | 236 |
| Alginate  | Hydrogels produced by combining<br>glutaraldehyde-stabilized gelatin<br>and natural polysaccharide alginate   | Lipase                                | Cross linking                                     | Stable and can be recycled for 10 and 20 times in aqueous and micellar media respectively without noticeably losing enzyme activity  | 237 |
| Alginate  | Microcapsules with an alginate-<br>chitosan shell   | β-Galactosidase                       | Entrapment  | (1) Ca <sup>2+</sup> alginate has a lesser loading efficiency than Ba <sup>2+</sup> alginate. (2) Compared to liquid core Ca <sup>2+</sup> alginate microcapsules and the unbound enzyme, solid core Ba <sup>2+</sup> alginate microcapsules increased | 238 |
| Alginate  | Ultrathin alginate/protamine/silica<br>(APSi) hybrid membranes in the<br>core-shell capsules  | Laccase                               | Entrapment  | the stability of the enzyme at 37 °C After 20 days, the stability of the encapsulated laccase increased by 67%. Improved storage, pH, and thermal stabilities  | 239 |
| Alginate  | Alginate-chitosan microcapsules<br>made using an internal gelation<br>and emulsification process to create<br>alginate beads  | Laccase                               | Not mentioned                                     | Increased loading effectiveness and stability. In the Alizarin Red dye decolorization test, the free and immobilised laccase alone both had very low decolorization efficiency   | 240 |
| Alginate gel  | Alginate-silica nanotubes (ALG-SiNTs) composite was created by incorporating silica nanotubes (SiNTs) into the alginate (ALG) gel and then encapsulating them with Ca <sup>2+</sup> through cross-linking | Yeast alcohol<br>dehydrogenase (YADH) | Encapsulation                                     | Improved storage and operational<br>stability with ALG-SiNT composite<br>showing ~50% lesser leaching<br>compared to only alginate matrix  | 241 |
| Alginate  | Biomaterials made from alginate gel<br>doped with double-walled carbon<br>nanotubes (DWCNT-ALG)   | LDH                                   | Adsorption,<br>encapsulation and<br>cross linking | Improved enzyme adherence with >61% reduction in leaching and 25 folds higher shelf-life than native enzyme  | 243 |
| Alginate  | Ca-alginate beads by entrapment   | Lipase                                | Entrapment  | 40% higher activity than the free<br>enzyme with 10 cycles of reusability<br>without any loss of activity  | 244 |
| Alginate  | Calcium alginate beads  | Protease                              | Entrapment  | Activity retained for a longer period<br>of time and reused for 3 cycles. Up<br>to 10 days of storage stability at 4 °C<br>was observed  | 245 |
| Alginate  | Calcium alginate gel by entrapment method   | Cellulase                             | Entrapment  | Increased stability with 5 °C raise in optimum temperature (55 to 60 °C) and pH tolerance in acidic regime   | 246 |
| Alginate  | Calcium-alginate beads made using the entrapment technique  | MnP                                   | Entrapment  | (1) >83% immobilization yield;<br>tremendous shift in the optimum<br>temperature from 35 to 60 °C and<br>pH (towards acidic range). (2)<br>>82–95% decolouration capacity<br>with Sandal-fix dyes  | 247 |
| Sodium salt of<br>alginic acid<br>from brown<br>algae | PVA-ALG beads   | Naringinase                           | Entrapment  | Residual activity retained 70% after<br>8 successive batches   | 248 |
| Alginate  | PVA with calcium alginate (Alg) beads   | Lipase                                | Entrapment  | Improved the thermal stability and reusability   | 249 |
| Alginate  | Alginate-glutaraldehyde beads made through covalent bonding   | Neutrase                              | Covalent bonding                                  | Immobilization yield was ~50% and bestowed 10 °C higher thermal tolerance than free enzyme (50 °C)   | 250 |
| Alginate  | $\rho\textsc{-Benzoquinone-activated}$ alginate beads   | Glucoamylase                          | Entrapment  | No change in optimum pH,<br>temperature and activity over 36<br>days and 30 cycles of reycling   | 251 |
| Alginate  | Calcium alginate-chitosan<br>microspheres (CACM)  | GOx                                   | Encapsulation and adsorption method               | Increased the enzymatic activity and storage stability was increased up to 2 month and retained activity of CACM-GOX   | 252 |
| Alginate  | Hybrid gel beads made of<br>chemically reduced graphene oxide<br>(CRGO) and alginate  | GOx                                   | Non-covalent<br>adsorption-<br>entrapment method  | Good recyclability, mechanical properties, and enhanced stability in a wide pH (4–6) and temperature (45–60 °C) range  | 253 |

Table 6 (Contd.)

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| Biomass  | Biomass derived material   | Used enzyme   | Immobilization<br>method                  | Result/performance study  | Ref. |
|----------|--|---|---|---|------|
| Alginate | Polymethyl methacrylate (PMMA-g-CS) nanoparticles were grafted onto chitosan encapsulated calcium alginate beads | α-СТ  | Covalent bonding                          | Higher retained activity at pH 9 for 24 h   | 254  |
| Alginate | polyvinyl alcohol/alginate and<br>polyethylene oxide/alginate<br>nanofibers by electrospinning<br>method         | Lipase  | Adsorption and cross linking method       | Enhanced the stability. Maintained 60% of their activities after 14 and 7 reuses and increased the stability  | 256  |
| Alginate | Nanoflower/alginate microbeads   | ALDC  | Entrapment                                | Restored 98% of activity with   | 257  |
| Alginate | Calcium alginate beads with glutaraldehyde activation  | Xylanase  | Entrapment and cross linking              | improving stability and recyclability<br>After 30 days at 4 °C, the enzyme<br>still exhibits 80% of its initial<br>activity and maintains recycling<br>efficiency up to five reaction cycles<br>with 37% retention activity | 258  |
| Alginate | Functionalized calcium alginate beads with chitosan coating  | Acrylamidase  | Cross linking and covalent immobilization | Enhanced pH, thermal, shelf, and<br>mechanical stability, and<br>maintained 80% activity after four<br>cycles   | 259  |
| Alginate | Activated alginate-montmorillonite (MMT) beads   | Pectinase   | Covalent binding                          | Showed greater activity, however the optimal pH dropped from 5.5to 5.0. The immobilised enzyme's initial activity was maintained at around 53% after 6 cycles of reuse  | 260  |
| Alginate | Glutaraldehyde is utilized to harden<br>hybrid alginate-chitosan beads   | Saccharomyces cerevisiae<br>alcohol dehydrogenase<br>(SCAD) | Cross linking                             | Improved thermal stability with<br>enhanced optimum temperature<br>(from 30 to 40 °C)   | 261  |
| Alginate | Boehmite/alginate hybrid beads   | YADH  | Encapsulation                             | After 67 hours of incubation,<br>encapsulated YADH can practically<br>approach "zero leaching" and retain<br>86.6% of its activity after 12 cycles  | 262  |
| Alginate | Entrapped BG crosslinked<br>glutaraldehyde in calcium alginate<br>particles                                      | β-Glucosidase   | Entrapment and crosslinking               | Under the optimum conditions,<br>more than 60% of enzyme activity<br>restored and cross linkage of<br>glutaraldehyde reduced leakage of<br>BG from the calcium alginate<br>particles  | 263  |
| Alginate | Polyvinyl alcohol-sodium alginate (PVA-SA) nanofibers by electrospun nanofiber                                   | Phytase   | Cross linking                             | Enhanced the catalytic activity with optimum pH shifting towards higher pH (from 5 to 6) and temperature from 45 to 55 °C   | 264  |
| Alginate | Entrapment of alginate beads   | Lipase  | Entrapment                                | High lipase activity in the acidic pH of 3 and 40 °C temperature  | 265  |
| Alginate | 3D transition metal cation-crosslinked alginate nanogels $(Mn^{2+}, Fe^{3+}, and Co^{2+})$                       | Urease  | Encapsulation                             | Fe <sup>3+</sup> -alginate crosslinked nanogels<br>demonstrated high enzyme activity,<br>efficient enzyme loading, and zeta<br>potential compared to Co <sup>2+</sup> and<br>Mn <sup>2+</sup>                               | 266  |
| Alginate | Cu-alginate beads  | Trametes Versicolor<br>laccase                              | Entrapment                                | Enhanced activity with >96%<br>degradation capacity towards<br>bisphenol at pH 5 and 30 °C in<br>60 min   | 268  |

abundant functional groups. <sup>261–268</sup> For instance, Kamaci *et al.* <sup>264</sup> observed enhanced catalytic activity of immobilized phytase into polyvinyl alcohol-sodium alginate (PVA-SA) electrospun nanofibers. Nanofibers were fabricated by mixing PVA and SA at a 80:20 ratio, voltage of 23 kV, and distance of 14 cm *via* electrospinning. Sharma *et al.* <sup>269</sup> prepared bio-based and low-cost hybrid alginate-protein cryogel beads as novel adsorbent materials for the purification of immunoglobulin (Ig)G from human serum (Fig. 24). Due to soft interaction between the protein and IgG, the stability and integrity of the antibody were retained after the desorption step. Similarly,



**Fig. 24** Use of hybrid alginate-protein cryogel beads to purify IgG (schematic). From ref. 269.

STRENGTHS **WEAKNESS** ·Energy-intensive biomass ·Renewable biomass Eco-friendly biocatalysts processing BDMFs with high stability Strong interaction between and reusability BDFMs and enzymes Limited use of BDMFs a industrial-scale **OPPORTUNITIES** THREATS ·Scalable process Irreversible damage to Scope for tandem protein's secondary biocatalysis structure Modification of Inefficient back extraction ncapsulated enzymes of native enzymes soft interactions

Fig. 25 SWOT analysis of BDFMs for biocatalysis.

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upcycling of marine chitin was improved with immobilized chitinolytic enzymes.<sup>270</sup>

### 8. Conclusions and future prospects

The use of enzyme-based biocatalysts at the industrial scale has garnered interest in recent decades. However, the use of enzymes is limited due to their low thermal stability, complicated reusability, self-cleavage, and aggregation under extreme processing conditions. Immobilization of enzymes on a biocompatible and renewable carrier is a promising method to overcome such drawbacks. BDFMs are promising carriers for enzyme immobilization due to their availability, biodegradability, biocompatibility, and low risk of environmental contamination. In this review, the main types of BDFMs were presented as promising carriers for enzymes to improve their stability and reusability. In addition, the immobilization of enzymes on various biomass-derived carriers (e.g., cellulose, lignin, SF, chitin, chitosan, and ALG) was surveyed. Although various approaches have been developed for BDFM fabrication, studies on the practical applications of these materials are in the early stages. Immobilization/adsorption/entrapment of various enzymes on BDFMs can improve the stability, activity, reusability, and recyclability of biocatalysts compared with those of their native state. In many cases, impressive results have been obtained whereas, in several other cases, the biocatalytic activity must be improved for commercial application. For selection of the best support materials and preparation of an efficient biocatalyst, all reaction conditions must be optimized in terms of monodispersity and surface chemistry. Moreover, the literature is based mainly on enzymes of a single type, which may not be suitable for industrial needs. Therefore, studies with multiple enzyme "cocktails" should be undertaken to lower the cost and increase the utility of biocatalysts.

Fig. 25 provides an overview of the analysis of the strength, weakness, opportunities and threats (SWOT) of BDFMs as carriers for enzymes. These challenges should be considered when selecting the biomass matrix and immobilization methods for direct industrial application. Cost-effective sources (e.g., non-commercial biomass, waste biomass, nonwood forest products, and biomass from unconventional resources) as support materials as well as controlled immobilization methods with soft interactions between enzymes and BDFMs must be selected. This review also provides an opportunity to understand the availability of different types of biomass as catalyst supports to develop efficient biocatalytic systems. The recyclability and reusability of enzymes is another crucial parameter that ensures the practicality of a biocatalytic process. We believe that this review will aid use of BDFMs for further studies with different enzymes under extreme conditions (e.g., high temperature, pH, denaturing substances). We attempted to collect all the information on the immobilization of enzymes on macro- or microscopic biomass-derived materials as carriers to improve the enzyme activity, stability, and recovery of biocatalysts. Therefore, this review aims to create economic value from various biomass and biocatalytic processes to strengthen the economy, with high scientific, technological, and societal impacts. This review also aids the use of biomaterials as supports for the development of immobilized biocatalysts for applications based on energy, environmental issues, and chemical syntheses. From the perspective of green and sustainable chemistry, this review offers a promising approach for effective evaluation of biomass.

#### **Abbreviations**

CRL

| BDFM       | Biomass-derived functional material  |
|------------|--------------------------------------|
| AA         | Alginic acid                         |
| MOF        | Metal-organic framework              |
| WOS        | Web of science                       |
| BGU        | β-Glucuronidase                      |
| CHM        | Cellulose hydrogel microsphere       |
| CNC        | Cellulose nanocrystal                |
| CNW        | Cellulose nanowhisker                |
| NFC        | Nanofibrillated cellulose            |
| MCC        | Microcrystalline cellulose           |
| CBD        | Cellulose-binding domain             |
| GOx        | Glucose oxidase                      |
| CMC        | Carboxymethyl cellulose              |
| IL         | Ionic liquid                         |
| PPL        | Porcine pancreatic lipase            |
| HNT        | Halloyite nanotube                   |
| MCM        | Cellulose porous microsphere         |
| PGA        | Penicillin G acylase                 |
| TLFCH      | Tendril-like functional carbon helix |
| DES        | Deep eutectic solvent                |
| Cyt c      | Cytochrome c                         |
| [Emim][Ac] | 1-Ethyl-3-methylimidazolium acetate  |

Candida rugosa lipase

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OPH Organophosphorus hydrolase
BTDE 1,4-Butanediol diglycidyl ether
EMBR Enzymatic membrane bioreactor

 $\begin{array}{ll} \alpha CF & \alpha\text{-Cellulose fiber} \\ \alpha\text{-CT} & \alpha\text{-Chymotrypsin} \end{array}$ 

PDDA Poly(diallyldimethylammonium chloride)

L-DOPA 3,4-Dihydroxyphenyl-L-alanine

LNP Lignin nanoparticle

SET-LRP Single electron transfer-living radical

polymerization

Horseradish peroxidase

SF Silk fibroin

HRP

ChOx Cholesterol oxidase CA Carbonic anhydrase **ASNase** L-Asparaginase CadA Lysine decarboxylase ChBD Chitin-binding domain **CHNW** Chitin nanowhisker AuNP Gold nanoparticle Silver nanoparticle AgNp CNF Chitin nanofibril

TMB 3,3',5,5'-Tetramethylbenzidine

GO Graphene oxide

RGO Reduced graphene oxide BSS Bamboo shoot shell LBL Layer-by-layer SGP Seal gastric protease

CAT Catalase

NaIO<sub>4</sub> Sodium periodate

TEMPO 2,2,6,6-Tetramethylpiperidine-1-oxyl radical

SNF Silk nanofiber

ONPG *o*-Nitrophenyl-β-p-galactopyranoside DWCNT Double-walled carbon nanotube

LDH Lactate dehydrogenase
PDMS Polydimethylsiloxane
PEO Polyethylene oxide

ALDC α-Acetolactate decarboxylase

EDC 1-Ethyl-3-(3-dimethylaminopropyl) carbodiimide

hydrochloride

NHS N-Hydroxysuccinimide

 $t_{1/2}$  Half life

PEG Polyethylene glycol
Ca-ALG Calcium-alginate
AAC Alginate-derived carbon

SiNT Silica nanotube TMOS Tetramethoxysilane

BCA Bovine carbonic anhydrase

SA Sodium alginate XA Xanthan-alginate

PVA-SA Polyvinyl alcohol-sodium alginate

MMT Montmorillonite

DMC Dialdehyde-modified cellulose nanocrystal MDC Magnetic dialdehyde cellulose nanoparticle

MCNC Magnetic chitin nanofiber composite

CS-Nps Chitosan nanoparticle ALG-SiNT Alginate-silica nanotube CMNP Chitosan magnetic nanoparticle

MnP Manganese peroxidase

#### Conflicts of interest

There are no conflicts to declare.

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