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On the use of modelling antagonistic enzymes to aid in temporal programming of pH and PVAborate gelation†

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Feedback through enzyme reactions creates new possibilities for the temporal programming of material properties in bioinspired applications, such as transient adhesives; however, there have been limited attempts to model such behavior. Here, we used two antagonistic enzymes, urease in watermelon seed powder and esterase, to temporally control the gelation of a poly(vinyl alcohol)-borate hydrogel in a onepot formulation. Urease produces base (ammonia), and esterase produces acid (acetic acid), generating a pH pulse, which was coupled with reversible complexation of PVA. For improved understanding of the pulse properties and gel lifetime, the pH profile was investigated by comparison of the experiments with kinetic simulations of the enzyme reactions and relevant equilibria. The model reproduced the general trends with the initial concentrations and was used to help identify conditions for pulse-like behaviour as the substrate concentrations were varied.

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Design, System, Application

Enzyme reactions with pH-driven feedback provide a convenient route for programming gelation and motion of polymers in applications such as soft robotics. There is a need to develop models to improve our understanding and control of such systems. Here, we investigated the transient formation of PVA-borate hydrogels driven by a combination of acid- and base-producing enzymes and compared experiments to kinetic simulations of the reactions and equilibria. The model reproduced the general trends and enabled a rapid search of parameters to find optimal conditions for the desired pulse-like

1. Introduction

There is increasing interest in the exploitation of feedback in the programming of smart materials, such as hydrogels or polymers that undergo conformational transitions (volume phase or gel-to-solution phase) in response to environmental signals.¹⁻³ Feedback arises through the pH when acidic or basic products catalyze the reaction; leading to rate acceleration and phenomena such as a pH clock (sharp switch in pH after a time lapse), pulse (transient increase/decrease in pH) oscillations.4,5 Transient changes in pH may be exploited as a

convenient method to control the lifetime of pH-sensitive hydrogels in bioinspired applications, such as temporary adhesives or soft robotics.^{6,7} Proposed mechanisms for temporary adhesion include secretion of an adhesive and deadhesive substance in aquatic organisms such as starfish⁸ and slime moulds achieve motion using periodic contraction of actin fibrils. These organisms have long been studied for their ability to perform seemingly complex spatial tasks, such as maze-solving.10 A better understanding of pH-driven feedback in soft matter is still required to achieve the level of selforganization obtained in biological systems.

Examples of using chemical feedback to trigger polymerization or chemomechanical responses mainly involve inorganic or toxic reactants. Hu et al. demonstrated a proof of concept of time-lapse curing of polymerization driven by a formaldehyde-sulfite pH clock.11 This reaction exhibits basecatalyzed feedback that results in an increase in pH after a controllable time lag, or clock time, and the sharp changes in state can be temporally programmed through the initial concentrations. The reaction has also been used for production

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of zeolites and chitosan particles. 12,13 Yoshida and collaborators achieved chemomechanical oscillations by coupling a pHsensitive hydrogel with the iodate-sulfite pH oscillating reaction in a flow reactor, and this was subsequently developed into a synthetic muscle. 14,15 The feedback in these systems involves acid autocatalysis and hydrogels with ionisable groups that change state according the pH of the environment. Novakovic and Isakova created a pH oscillating chitosan gel for pulsatile drug release using a palladium-catalyzed carbonylation reaction.¹⁶ In closed systems, only the Belousov-Zhabotinsky (BZ) reaction, which is autocatalytic in bromous acid, has been exploited for periodic changes. A self-oscillating gel was obtained with a ruthenium catalyst covalently bonded to the gel matrix and periodic polymerization was observed using the BZ reaction with acrylonitrile. 17,18 The major disadvantage of the BZ reaction, as well as most pH oscillators, is the harsh oxidizing environment.

Enzymes offer sustainable and benign approaches to controlling material formation in a closed system. 19,20 Compared to traditional methods that rely on harsh chemicals, enzymatic approaches can be coupled with sensitive biomolecules and bioactive components, expanding the potential for diverse applications including pacemakers, drug delivery systems, selfhealing polymers, self-walking actuators and transducers. 21,22 Only a small number of enzymes with feedback have been used to date for materials applications. Siegel et al. used glucose oxidase to create chemomechanical oscillations in a hydrogel system. 23,24 Acid production from glucose modulated the swelling of a hydrogel and periodically delivered a hormone across a membrane. A trypsin-based system was used to drive periodic assembly of complex coacervates.²⁵ Urease catalyzes the formation of ammonia from urea and has been widely used to control the timing of base-driven gelation, supramolecular polymerization and peptide self-assembly.^{7,19} Spatial pH patterns have also been produced, for example, the base produced by the reaction was used to catalyze a thiol-acrylate Michael addition reaction resulting in propagating hydrogel fronts and urease was combined with glucose oxidase to generate spatial gradients in pH in a hydrogel that were perpetuated in the presence of nanoparticles. 26,27 For a more complex temporal response, the coupling of pH clocks from opposing enzyme reactions with pHresponsive DNA hydrogels was demonstrated by Heinen et al.²⁸ The system utilized urease and esterase, which catalyzed the hydrolysis of ethyl acetate into acetic acid. Together, these enzymatic reactions generated a pH pulse, characterized by an initial increase followed by a decrease in pH, and the lifetime of the DNA gel was controlled.

There have been few attempts to model enzyme reactions with feedback for the temporal control of hydrogel properties. Mostly these focused on glucose oxidase-based reactions coupled to changes in hydrogel permeability, however a model was proposed for antagonistic enzymes involving bond formation and destruction in which it was demonstrated that transient gelation could occur. 20,23 Models have been developed for the feedbackdriven increase or decrease in pH with the urease or esterase reactions, but questions remain regarding their ability to predict

pH pulses in a multiphase system. Therefore, we combined experiments with modelling to show that these enzymes can be used to temporally control the gelation of polyvinyl alcohol (PVA) with borate. PVA-borate gels are non-toxic, biodegradable and inexpensive hydrogels that undergo pH responsive, reversible transitions that may be exploited in applications. 29-31 Formulations have been developed for topical drug release, although in recent years some concerns over boric acid toxicity have been raised. 32,33 Other applications include cleaning of painted surfaces in art conservation and PVA-borate gels are also widely used as model systems for studying physical gelation properties.34,35 Inorganic reactions have also been used for control of PVA gelation including autocatalytic iodate systems to form PVA-iodine gels, where it was demonstrated that the mechanical properties of the gel could be tuned through the changes in the reaction time. 36,37

Herein, we obtained transient gelation of PVA-borate in a onepot formulation containing the urease and esterase enzymes, buffer, boric acid and PVA solution. In order to determine the conditions for a pH pulse and transient gelation, a kinetic model was developed incorporating the enzyme reactions and equilibria with data obtained from the literature and our previous work.³⁸ The model reproduced the general trends with initial concentrations and was used to find optimal substrate concentrations for a pH pulse. However, a quantitative match between experiments and simulations was lacking in some cases, suggesting additional factors are required to fully explain the behavior.

2. Experimental section

2.1. Materials

Sodium acetate (3 M, pH 5.2) was purchased from VWR Chemicals. Poly(vinyl alcohol) (LMW, $M_{\rm w}$ = 31 000–50 000, 98– 99% hydrolysed), boric acid (ACS reagent, 99.5%), acetic acid (glacial), urea and ethyl acetate ($\rho = 0.902 \text{ g ml}^{-1}$) were purchased from Sigma Aldrich. The enzyme esterase was from porcine liver (Sigma, E3019-100KU) with an activity of 18 U mg⁻¹. All chemicals were used as purchased without further purification.

The watermelon seed powder (WMSP) containing the enzyme urease was prepared as described in our earlier work.³⁹ To summarize, Jubilee improved seeds were obtained from Eden brothers (Arden, USA) and were ground to a fine powder in a flour mill, ensuring that the mixture doesn't overheat. Acetone, in a ratio 2:1 of acetone to seed powder by volume, was used to delipidate the seeds and this mixture was left overnight. The seed mixture was filtered, and the cake washed again with acetone. The cake was allowed to dry out in a fume hood overnight. The activity of the urease in the batch of watermelon seed powder used in experiments was determined using Nessler's reagent and was 347.8 mg NH₃ per g WMSP per 5 minutes or 4084 U g⁻¹.40

2.2. Production of the PVA-borate hydrogels

Stock solutions were prepared of poly(vinyl alcohol) (PVA, 4-10 wt%), boric acid (5 wt%, 0.8 M), urea (1 or 0.5 g ml⁻¹

prepared by ultrasonication) and sodium acetate/acetic acid buffer (pH 4.05). The acetate buffer solution was prepared by adding 7.5 ml of acetic acid (glacial. 17.5 M) to 20 ml of 3 M sodium acetate and then acid added dropwise to obtain a final pH of 4.05. The buffer concentration was [acetate] + [acetic acid] = 6.9 M.

For the formation of PVA-borate hydrogels, typically 10 ml of PVA solution was added to a 25 ml round bottom flask and then boric acid solution (5 wt%) was added dropwise with swirling. The watermelon seed powder (WMSP) was added to the flask, and the solution was vortexed for 5 minutes. The esterase powder was then added to the flask, and the mixture vortexed again for 5 minutes. Then the buffer solution was added with a micropipette. The flask was placed on an Anzeser magnetic stirrer hotplate. A Vernier pH probe was inserted into the mixture, and the pH was monitored until it stabilized. The reaction was initiated by simultaneous addition of the urea solution and ethyl acetate and the mixture was constantly stirred using a magnetic flea at a stirring rate of 800 rpm. Complete gelation was recorded when the motion of the flea stopped. The flask was covered with foil while the reaction proceeded to reduce loss of ammonia. Experiments were performed at room temperature (25 °C).

Unless otherwise stated, the mixture composition for a pulse was: 10 mL PVA (4 wt%) solution, 50 µL of boric acid solution (5 wt%), 0.45 g of WMSP, 0.15 g of esterase, 100 µL acetate buffer (6.9 M), 400 µL of urea (1 g ml⁻¹), and 1050 μL of ethyl acetate. Thus, the standard concentrations were: urease activity = 184 U ml⁻¹, esterase activity = 270 U ml⁻¹, $[PVA] = 40 \text{ g L}^{-1}$, [boric acid] = 4 mM, [acetate buffer] =0.0693 M, [urea] = 0.667 M and [ethyl acetate] = 1.08 M. The effect of buffer concentration on the pH pulse was investigated by varying the volume of acetate buffer from 100 to 800 µL in separate experimental runs. The effect of urease concentration on the pH pulse was investigated by varying the mass of the watermelon seed powder from 0.15 g to 0.90 g and the effect of urea concentration was investigated by varying the volume of urea solution added from 100 to 800 µL. Variations in esterase, PVA and boric acid concentrations were also investigated in pulses experiments. Repeated were obtained simultaneous additions of the urea solution and ethyl acetate after each pulse. The viscosity of the mixture in time was determined using a Brookfield DV-II+ viscometer with a LV2-62 spindle at 50 RPM with a 10% PVA solution and the other concentrations as above.

3. Modelling of the coupled ureaseesterase reaction

The system was modelled using a set of coupled ordinary differential equations (ODEs) that describe the rate of change of species within the solution. The main processes in the model were (3.1) the urease-catalyzed reaction, (3.2) the esterase-catalyzed reaction, and (3.3) the equilibria that govern the pH (3.3). We used a bell-shaped dependence of the rate on pH for urease (in WMSP) and esterase with a maximum rate at pH \sim 8.

3.1. Urease-catalyzed reaction

Urease-catalyzed hydrolysis of urea yields ammonia and carbon dioxide:

$$CO(NH_2)_2 + H_2O \stackrel{urease}{\rightarrow} NH_3 + CO_2$$
 (1)

The rate of the enzyme-catalyzed reaction was given by a modified Michaelis-Menten equation (for simplicity acid is included as H⁺ rather than H₃O⁺):^{41,42}

$$\nu_{0} = \frac{k_{1a}[E]_{T}[U]}{\left(K_{M}\left(1 + \frac{[BA]}{K_{I}}\right) + [U]\left(1 + \frac{[U]}{K_{S}}\right)\right)\left(1 + \frac{K_{es2}}{[H^{+}]} + \frac{[H^{+}]}{K_{es1}}\right)\left(1 + \frac{[NH_{4}^{+}]}{K_{P}}\right)}$$
(2)

where k_1 is the rate constant for irreversible decomposition of the enzyme-substrate complex into products, [E_T] is the concentration of enzyme, [U] is the concentration of urea, K_M is the Michaelis constant, [H+] is the concentration of acid, and K_{es2} and K_{es1} are the protonation equilibria of the substrateenzyme complex that give rise to the bell-shaped rate-pH curve (see ESI†). Substrate and product inhibition terms were included: K_S = equilibrium constant for uncompetitive substrate inhibition and K_P = equilibrium constant for non-competitive product inhibition; as well as competitive inhibition from boric acid (BA) with constant $K_{\rm I}$ (SI).

The turnover number $(k_{\text{cat}} \text{ (s}^{-1}))$ of urease varies depending on the source, purity and conditions of the assay.41 In our previous work with urease type III from jack bean (Sigma), we defined $k_{1a} = k_{cat}/p/Mr \times (1000 \text{ ml dm}^{-3})$, where p = specific activity, Mr is molecular mass and with $[E]_T$ given as activity in U ml⁻¹ so that the product $k_1[E]_T$ gives the maximum rate, V_{max} in M s⁻¹.³⁸ There is limited experimental data for urease contained in WMSP (which is not pure), so we took a value of $k_{\text{cat}} = 5500 \text{ s}^{-1}$ in line with our previous work and used $p = 3750 \text{ U mg}^{-1}$, Mr = 470 kDa and $K_{\rm M}$ = 8 mM for urease extracted from WMSP to give a value of $k_{1a} = 3.12 \times 10^{-6} \text{ M U}^{-1} \text{ ml s}^{-1}$. For WMS urease, the pH optimum was ~8,39,43 so here we used values of K_{es1} and K_{es2} to give that optimum (Fig. S1†), and the values of K_s and K_I within ranges given the literature. We adjusted the value K_p to give a better match to experiments (see discussion). The rate constants are listed in Table 1.

3.2. Esterase-catalyzed reaction

Esterase-catalyzed hydrolysis of ethyl acetate yields acetic acid and ethanol:

Table 1 Rate constants (20 °C) and enzyme constants for the urease-esterase pH pulse

Equilibria rate constants	$k_2 \text{ s}^{-1}$ 7.3×10^5 $k_6 \text{ s}^{-1}$ 0.037	$k_{2r} \text{ M}^{-1} \text{ s}^{-1}$ 4.2×10^{10} $k_{6r} \text{ M}^{-1} \text{ s}^{-1}$ 7.9×10^{4}	$k_3 \text{ s}^{-1}$ 1.25×10^{-6} $k_7 \text{ s}^{-1}$ 2.8	$k_{3r} \text{ M}^{-1} \text{ s}^{-1}$ 1×10^{10} $k_{7r} \text{ M}^{-1} \text{ s}^{-1}$ 5×10^{10}	$k_4 \text{ M s}^{-1}$ 1×10^{-4} $k_8 \text{ s}^{-1}$ 5.794	$k_{4r} \text{ M}^{-1} \text{ s}^{-1}$ 1×10^{10} $k_{8r} \text{ M}^{-1} \text{ s}^{-1}$ 1×10^{-10}	$k_5 \text{ s}^{-1}$ 24 $k_9 \text{ M}^{-1} \text{ s}^{-1}$ 200	$k_{5r} \text{ M}^{-1} \text{ s}^{-1}$ 4.3×10^{10} $k_{9r} \text{ s}^{-1}$
Urease enzyme constants	$k_{1a} \text{ M U}^{-1} \text{ ml s}^{-1}$ 3.12 × 10 ⁻⁶		$K_{\text{Ma}} \text{ M}$ 8×10^{-3}	$K_{es1a} \\ 8 \times 10^{-7}$	K_{es2a} 2×10^{-11}	<i>K</i> _s M 3	<i>К</i> _р М	$K_{ m I}~{ m M}$ $1 imes10^{-4}$
Esterase enzyme constants	$k_{1b} \text{ M U}^{-1} \text{ ml s}^{-1}$ 1.17 × 10 ⁻⁶		$K_{\rm Mb} M$ 3×10^{-3}	$\begin{array}{c} K_{\rm es1b} \\ 2 \times 10^{-6} \end{array}$	$K_{es2b} \\ 2 \times 10^{-11}$			

The rate of the enzyme-catalyzed reaction was given by a modified Michaelis-Menten equation (for simplicity, acid is included as H⁺ rather than H₃O⁺):

$$\nu_{0} = \frac{k_{1b}[E]_{T}[EA]}{(K_{M} + [EA])\left(1 + \frac{K_{es2}}{|H^{+}|} + \frac{[H^{+}]}{K_{es1}}\right)}$$
(4)

where k_{1b} is the rate constant for irreversible decomposition of the enzyme-substrate complex into products (s⁻¹), [E]_T is the concentration of enzyme, [EA] is the concentration of ethyl acetate, $K_{\rm M}$ is the Michaelis constant, $[{\rm H}^{+}]$ is the concentration of acid, and K_{es1} and K_{es2} are the protonation equilibria of the substrate-enzyme complex (see ESI†).

The form of the rate equation for esterase depends on the source and heterogeneity of enzyme, and nature of the substrate. There was limited experimental data on the rate for pig's liver esterase with ethyl acetate, however, early work suggested a small deviation from Michaelis-Menten kinetics and the value of $k_{\text{cat}} = 3.55 \text{ s}^{-1}$ and $K_{\text{M}} = 1 \text{ mM}$ (pH 8, 25 °C).44 With the molecular weight of 168 kDa and specific activity of 18 U mg⁻¹ for esterase purchased from Sigma, we took a value of $k_{1b} = 1.17 \times 10^{-6} \text{ M U}^{-1} \text{ ml s}^{-1}$ and with $[E]_T$ recorded in U ml⁻¹ so that the product $k_1[E]_T$ gives the maximum rate, V_{max} in M s⁻¹. The pH-rate curve in earlier work also varied with the enzyme source and substrate and the maximum rate was between 6.5 and 8, with a high activity maintained up to pH = 10, so here we used values of Kes1 and Kes2 to give that broad optimum (Fig. S1†). 28,45,46

3.3. pH equilibria

When both the urease and esterase reactions are considered together, the pH is determined by the following reversible reactions:

The acid equilibria rate constants are well established. 47-49 The equilibrium constant, K, for the boric acid-PVA complex was taken from the literature⁵⁰ where [diol] is the concentration of the diol subunit of PVA, calculated from the mass of PVA in experiments and $Mr = 88 \text{ g mol}^{-1}$. We used only the monodiol complexation reaction here, assuming the didiol complex is not significant at these borate concentrations.

3.4. Model equations and parameters

Reactions 1 and 3 and R2-R9 produce 17 coupled rate equations, including the pH determining equilibria and the two enzyme rates which are of a modified Michaelis-Menten form. The values of all the rate constants taken in this work are shown in Table 1. The rate equations were solved using XPPAUT with integration method CVODE (model file included in ESI†).51 The phase diagram was produced by incrementally increasing the values of one substrate with the other fixed and recording the data for a total of 1000 minutes to determine the magnitude of the pulse from the difference between the maximum pH and final pH at 1000 minutes.

A typical mixture composition for the simulations of the pH pulse was [urease] = 184 U ml⁻¹, [esterase] = 267 U ml⁻¹, [acetate buffer] = 0.0693 M, [ethyl acetate] = 1.08 M, [urea] = 0.667 M, [boric acid] = 4 mM, [diol] = 0.45 M.

4. Results

A schematic representation of the proposed system can be seen in Fig. 1. The coupling of the urease and esterase reactions leads to a pH pulse, i.e., a rise then fall in pH, for certain initial concentrations (Fig. 1a). The urease reaction is slow initially as the solution is at low pH. The ammonia produced from hydrolysis of urea results in an increase in pH and a shift of the boric acid equilibrium to borate anion which is used to cross-link the PVA gel (Fig. 1b). However,

R2	$CH_3COOH = H^+ + CH_3COO^-$	$pK_a = 4.79$	Rate = $k_2[CH_3COOH] - k_{2r}[CH_3COO^-][H^+]$
R3	$CH_3CH_2OH \Rightarrow H^+ + CH_3CH_2O^-$	$pK_a = 15.9$	Rate = $k_3[CH_3CH_2OH] - k_{3r}[CH_3CH_2O^-][H^+]$
R4	$H_2O \rightleftharpoons H^+ + OH^-$	$pK_a = 14$	Rate = $k_4 - k_{4r}[OH^{-}][H^{+}]$
R5	$NH_4^+ \rightleftharpoons NH_3 + H^+$	$pK_a = 9.25$	Rate = $k_5[NH_4^+] - k_{5r}[NH_3][H^+]$
R6	$CO_2 + H_2O \rightleftharpoons H^+ + HCO_3^-$	$pK_a = 6.35$	Rate = $k_6[CO_2] - k_{6r}[HC_3][H^+]$
R7	$HCO_3^- \rightleftharpoons H^+ + CO_3^{2-}$	$pK_a = 10.25$	Rate = $k_7[HCO_3^-] - k_{7r}[CO_3^{2-}][H^+]$
R8	$B(OH)_3 + H_2O \Rightarrow B(OH)_4^- + H^+$	$pK_a = 9.24$	Rate = $k_8[B(OH)_3] - k_{8r}[B(OH)_4^-][H^+]$
R9	$B(OH)_4^- + diol = (B(OH)_4 - diol)$	K = 200	Rate = $k_9[B(OH)_3][diol] - k_{9r}[(B(OH)_4 - diol)]$

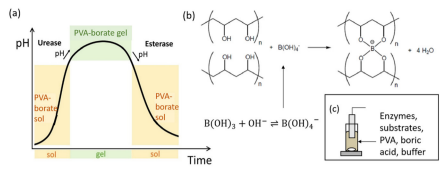


Fig. 1 (a) Schematic of time-controlled gelation using a pH pulse generated by kinetically balancing urease and esterase enzymatic reactions. There is a transient alkaline pH where the PVA-borax solution gels. (b) The shift of the boric acid equilibrium to borate with an increase in pH and crosslinking of the gel. (c) One-pot formulation employed in experiments.

the rate of the esterase reaction also increases, producing acetic acid, which then lowers the pH, and the de-gelation of PVA-borate occurs. For autonomous control, we employed a one-pot formulation containing enzymes and substrates, PVA, boric acid and buffer (Fig. 1c). Watermelon seed powder (WMSP) served as a cheap, easily accessible source of urease in this system.

A typical pH pulse during the experiment is shown in Fig. 2(a). The maximum pH and the steady state pH at 1000 minutes (pH_{final}), are indicated on the plot as well as the pulse magnitude, defined here as the difference between pH_{max} and pH_{final}. In general, with sufficient boric acid and PVA, for a pH_{max} greater than \sim 7.5 gelation of the mixture was observed, as illustrated in Fig. 2(b). If the final pH fell below \sim 6, then de-gelation of the mixture was obtained. In theory, gelation is completely reversible, and it was found that with repeated addition of the substrates, urea and ethyl acetate, multiple pulses could be obtained (Fig. 2(c)). However, the minimum pH failed to fall below 7 after several additions, likely because of deactivation of the esterase enzyme, and de-gelation was not obtained on this timescale. The transient increase in viscosity observed during a pH pulse was followed using a viscometer and is shown in Fig. 2(d) with max value of 270 cP. A range of states at pH_{max} was obtained, from a highly viscous solution to a gel, as the concentrations of PVA and boric acid were increased. The properties of borax-hydrogels are known to depend on the concentration of the PVA and boric acid, as well as the pH, ^{29,35} and addition of co-solvents such as propanol also influence the rheological properties.34

In this paper we focus on the kinetics of the pH changes. The characteristic features of the pH pulse, pH_{max} and

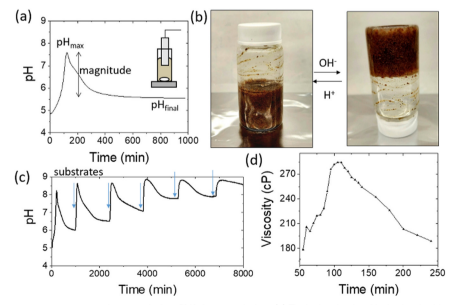


Fig. 2 Illustration of the esterase-urease reaction coupled to PVA-borax gelation. (a) Typical pulse in pH obtained with maximum pH and pH final (1000 minutes) indicated and urease in watermelon seed powder (0.045 g ml⁻¹; 184 U ml⁻¹), esterase (0.015 g ml⁻¹, 270 U ml⁻¹), [PVA] = 4 wt%, [boric acid] = 4 mM, [acetate buffer] = 0.0693 M, [urea] = 0.667 M and [ethyl acetate] = 1.08 M. (b) Formulation in vial, illustrating reversible gelation with increase and decrease in pH (c) repeated addition of substrates, urea and ethyl acetate (indicated by blue arrow) (d) viscosity measurements in time obtained with a Brookfield DV-II+ viscometer.

pH_{final}, and hence the temporal properties of the gel can be controlled by varying the initial concentrations and the kinetic balance between the base producing and acid producing reactions. We determined the general trends and compared them to kinetic simulations for different experimental conditions: varying the boric acid or buffer concentration, varying the substrate concentration and varying the watermelon seed powder (WMSP) or esterase concentration. The comparison between the experimental observations and the simulated results allows us to better understand the mechanism and the potential additional factors that may influence the pH-time pulse.

4.1. Effect of boric acid and acetate buffer concentration on the pH pulse

Pulses obtained in experiments with and without boric acid are shown in Fig. 3(a). The addition of boric acid decreases the maximum pH and increases the peak time. These results are qualitatively reproduced in the simulations with the inclusion of inhibition of urease by boric acid (Fig. 3(b)). Boric acid is removed by complexation with PVA during the pulse (Fig. 3(c)), and the pH_{max} is higher compared to reaction without PVA (Fig. 3(c) inset). In the simulation, urea is also completely removed during a pulse and the rate of removal of both substrates is observed to increase during the reaction, indicative of feedback (Fig. 3(d) inset). Feedback arises in the urease reaction as a result of the increase in rate of the enzyme reaction with an increase in pH.³⁸

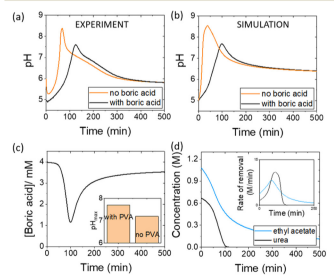


Fig. 3 Effect of boric acid on the urease-esterase pH pulse. (a) Experimental pH-time traces, prepared using urease in watermelon seed powder (0.045 g ml⁻¹), esterase (0.015 g ml⁻¹), [PVA] = 4 wt%, [boric acid] = 4 mM or 0 mM, [urea] = 0.667 M and [ethyl acetate] = 1.08 M. (b)-(d) Simulations of pulse with [urease] = 182 U ml⁻¹, [esterase] = 270 U ml⁻¹, [ethyl acetate] = 1.08 M, [urea] = 0.667 M, [boric acid] = 4 mM: (b) pHtime trace. (c) Concentration of boric acid in time and inset shows the pH_{max} with and without PVA, (d) concentration of substrates in time and inset shows their rate of change.

The acetic acid buffer can be used to increase the peak time (time to pH_{max}). The watermelon seed powder raised the initial pH of the solution to ~6.5 in the absence of buffer and the time to pH_{max} was 27 minutes (Fig. 4(a)i). With a buffer concentration = 0.069 M, the pH increased from \sim 4.5 to pH_{max} \sim 7.5 after 100 minutes. However, the magnitude of the pulse, defined here as pH_{max}-pH_{final}, decreased with the increase in peak time (Fig. 4(a)ii). The buffer kept the pH low, suppressing the urease reaction and reducing the pH maximum (Fig. 4(a)iii). The simulated profile also showed a pronounced pH pulse for [Ac buffer] < 0.1 M (Fig. 4(b)i), and the same general trends were obtained as in the experiments with increasing acetate buffer concentration (Fig. 3(b)ii and iii).

4.2. Effect of varying enzyme concentration on the pH pulse

In experiments with increasing mass of WMSP (Fig. 5), and hence urease activity, the magnitude of the pulse increased along with a decrease in the peak time (Fig. 5(a)ii). The faster rate of production of ammonia by urease raised the pH_{max} higher before the esterase could take effect (Fig. 5(a)iii). These general trends were qualitatively reproduced in the simulations with increasing activity of urease (Fig. 5(b)), however, the effect of urease on the maximum pH was more pronounced than in the experiments. Increasing the concentration of esterase had the reverse effect to urease, which was also reproduced qualitatively in the model (Fig. S2†).

4.3. Effect of varying substrate concentration on the pH pulse

The simulations can be used to rapidly map out parameter space to find the optimal conditions for a pH pulse. A phase diagram of pulse magnitude, is shown in Fig. 6(a) as a function of the two substrates, urea and ethyl acetate. The colour map indicates the pH change, in which dark blue shows the largest magnitude of pulse. For a given value of ethyl acetate concentration, there is an optimal range of urea concentrations for which pulse behaviour is obtained, and this expands as the concentration of ethyl acetate is increased.

The magnitude of the pulse increased then decreased with an increase in urea concentration and with [EA] = 1.08 M, the maximum magnitude was obtained at [U] = 0.67 M (Fig. 6(b)). The concentration of urea had a relatively small effect on the peak times compared to other parameters. The relationship between pulse magnitude and peak time is shown in Fig. 6(c) for two different concentrations of ethyl acetate. A decrease in [EA] to 0.6 M resulted in a larger maximum magnitude, but the peak times were also reduced.

Illustrative experiments at two different EA concentrations are shown in Fig. 6(d). As the [EA] was increased, the final pH was lower, as expected from the faster rate of production of acid. With [EA] = 1.08 M in experiments, a broad range of [U] gave pulse-like behaviour and the maximum pulse magnitude was obtained for [U] = 0.67 M, in good agreement with the simulations. No pulse-like behaviour was observed for [U] < 0.5 M and the process was dominated by esterase.

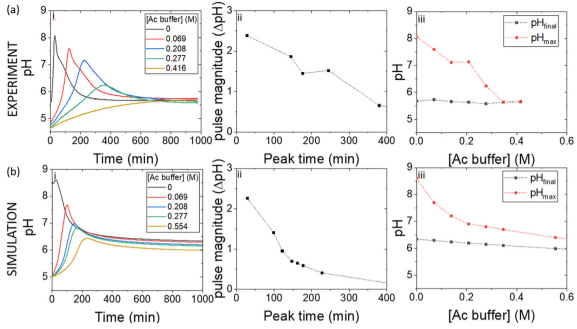


Fig. 4 Effect of the acetate buffer concentration on the urease-esterase pH pulse. (a) Experiments with urease in watermelon seed powder (0.045 g m $^{-1}$), esterase (0.015 g m $^{-1}$), [PVA] = 4 wt%, [boric acid] = 4 mM, [urea] = 0.667 M and (b) simulations with [urease] = 182 U m $^{-1}$, [esterase] = 270 U ml $^{-1}$, [ethyl acetate] = 1.08 M, [urea] = 0.667 M, [boric acid] = 4 mM. Effect of acetate on: (i) pH-time trace (ii) pulse magnitude (pH_{max} pH_{final}) vs. peak time and (iii) the maximum and final pH (T = 1000 min).

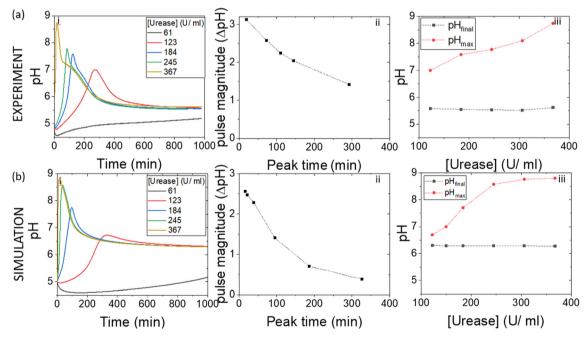


Fig. 5 Effect of the initial urease concentration on the urease-esterase pH pulse. (a) Experimental pH-time traces with: esterase (0.015 g ml⁻¹), [PVA] = 4%, [boric acid] = 4 mM, [acetate buffer] = 0.0693 M, [urea] = 0.667 M and [ethyl acetate] = 1.18 M. (b) Simulations of the pH pulse with [esterase] = 267 U ml⁻¹, [ethyl acetate] = 1.08 M, [acetate buffer] = 0.0693 M, [urea] = 0.667 M, [boric acid] = 4 mM. Effect of urease on (i) pH-time trace (ii) pulse magnitude vs. peak time, (iii) the pH maximum and final pH (T = 1000 min).

5. Discussion

It has been established in previous work that combining a base producing enzyme reaction and acid producing enzyme reaction can result in a pH pulse for temporal control of physical processes such as gelation.²⁸ Herein, we investigated the widely-used urease and esterase reactions coupled to PVA-borate gelation in order to determine how simulations

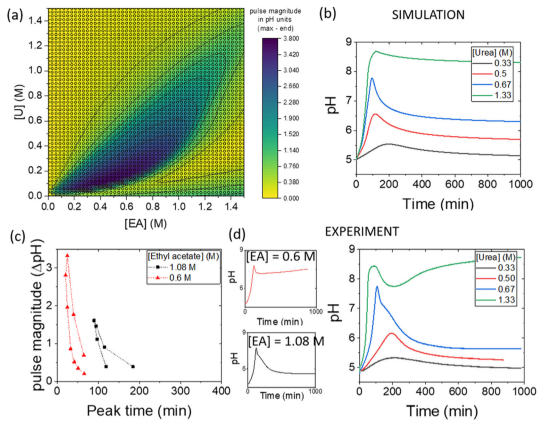


Fig. 6 (a) Simulated phase diagram showing pulse magnitude (colour bar) as a function of substrate concentrations, urea (U) and ethyl acetate (EA), with [urease] = 182 U ml⁻¹, [esterase] = 267 U ml⁻¹, [acetate buffer] = 0.0693 M, [boric acid] = 4 mM. (b) Simulated pH-time traces with variations in urea and [EA] = 1.08 M. (c) Change in pulse magnitude with peak time for two different concentrations of EA. (d) Experimental pHtime traces comparing [EA] = 0.6 M and 1.08 M and with variations in urea with [EA] = 1.08 M and urease in watermelon seed powder (0.045 g), esterase (0.015 g), [PVA] = 4 wt%, [boric acid] = 4 mM, [acetate buffer] = 0.0693 M.

may be used to aid in finding optimal conditions for a pulse. The rate of change of pH will influence both the properties and lifetime of the gel state³⁷ and the nature of the optimum will depend on the application. We focused on how the pHtime profile depends on the initial composition of the reaction mixture with comparison of the characteristic features of the pulse magnitude (pH_{max} - pH_{final}) and peak time in the experiments and simulations.

Overall, the model captured the general trends of pulse magnitude and peak time with initial concentrations. The simulations illustrated the switch from the urease rate to the esterase rate dominating as the reaction proceeds and the rate acceleration associated with the increase in pH (Fig. 3(d)). An increase in urease concentration (or decrease in esterase) resulted in a higher pH_{max} and there was an optimal substrate concentration for the largest pulse magnitude (Fig 6(a)). In general, there is a trade-off between pulse magnitude and peak time, as an increase in the magnitude arises from faster urease reaction relative to esterase, and hence shorter time to pH_{max}.

The model was less effective at producing a quantitative match to the experimental data and the results were found to be particularly sensitive to some of the enzyme parameters. The effect of halving or doubling k_1 or K_{es1} (the low pH binding site) on peak time and pH_{max} are shown in Table 2. Changing these parameters for urease results in ~60-500% change in the peak time and $\sim 10-15\%$ change in pH_{max}. We found that changing $K_{\rm M}$ or $K_{\rm es2}$ had no effect on the pH-time profile under the conditions explored here. We mainly used values from the literature for the enzyme constants, with some minor adjustments to improve the fit to the experiments. However, we note that for urease contained in

Table 2 Effect of changing enzyme constants k_1 and K_{es1} on peak time and pH_{max}

Esterase	% change in peak time	% change in pH_{max}	Urease	% change in T	% change in pH _{max}
$0.5 \times K_{\text{es1b}}$	-54	9.1	$0.5 \times K_{\mathrm{es1a}}$	477	-14.3
$2 \times K_{es1b}$	115	-11.7	$2 \times K_{es1a}$	-61	9.1
$0.5 \times k_{1b}$	-65.7	14.3	$0.5 \times k_{1a}$	588	-15.6
$2 \times k_{1b}$	250	-15.6	$2 \times k_{1a}$	-83.5	14.3

WMSP, and for esterase with ethyl acetate specifically, there was limited data available. 43,52,53 For urease from jack bean, a more complex dependence of the rate on pH has been proposed.42 Additionally, features associated heterogeneous forms of the esterase enzyme were not taken into account.44 Earlier work suggests that there are multiple binding sites possible, with activities depending on the substrate chain length and concentration, and the different isoforms have different optimal pH's.53 A more complex expression rather than the modified MM used here would likely improve the fit to the data; however this would result in more fitting parameters and hence we chose the simpler form here.

The importance of including the inhibition of urease by boric acid was also demonstrated, with an inhibition constant, $K_{\rm I}$, similar to that of jack bean urease giving a reasonable match between simulations and experiments.42 The substrate inhibition term K_s was found to play a small role in the dynamics, however the product inhibition $K_{\rm p}$ was found to completely suppress activity with the typical literature value of 0.002 used in our earlier work.³⁸ It is possible that product inhibition is different for urease contained in WMSP; this requires further investigation. However, the value of $K_{\rm I}$ was also previously determined for jack bean urease under conditions of much lower substrate, and hence product, compared to the conditions used here.41

Factors that were not included in the model include mixing effects caused by inhomogeneity in the distribution of watermelon seed powder and the influence of the increase in viscosity on the reaction rate. In experiments with no boric acid, the increase in viscosity with PVA from 1-12% was found to have a small effect on the peak time and value of pH_{max} (Fig. S2(a)†). However, with boric acid included, the change in peak time was greater, suggesting the dynamic change in viscosity may affect the rate of the urease reaction relative to the esterase reaction (Fig. S2(b)†). The diffusive restriction in the watermelon seed particles may also have influenced the effective enzyme rate compared to the simulations. For a better quantitative match to experiments, these features may be incorporated into the model in the future.

6. Conclusions

In conclusion, we have demonstrated the use of antagonistic enzymes, urease in WMSP and esterase, to obtain a pH pulse and coupled it to reversible gelation of PVA in a one-pot formulation. Our findings highlight the ability to modulate the pH-time pulse profile by manipulating experimentally controllable parameters. By varying the concentrations of substrate, buffer, and WMSP, we observed distinct changes in the pH dynamics. Repeated addition of substrates gave rise to multiple pulses, demonstrating the possibility of multiple gel cycles. The simulations reproduced the experimental trends qualitatively but there was a quantitative shift in the behavior compared to the experiments. These findings suggest that additional factors may be important, such as the more complex dependence of enzyme rate on pH or influence of increased viscosity and mixing effects on enzyme rate. Further investigation is needed to understand these issues and refine the model to better capture the experimental trends and to obtain control of the temporal programming of pH and gel properties. Overall, these results contribute to our understanding of coupled enzyme-catalyzed reactions and aid in the optimization of enzyme systems for various material applications.

Author contributions

JAP conceived of the idea, NB provided lab supervision and performed experiments; MM, IH, FS performed experiments; JB performed the viscosity measurements; AL and AFT wrote the model script and performed the simulations; AL and AFT wrote the original manuscript and all authors contributed to final version of the paper.

Conflicts of interest

There are no conflicts to declare.

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