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High Pressure Intensification of Cassava Resistant Starch (RS3) Yields

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Abstract

Cassava starch, typically, has resistant starch type 3 (RS3) content of 2.4%. This paper shows that the RS3 yields can be substantially enhanced by debranching cassava starch using pullulanase followed by high pressure or cyclic high-pressure annealing. RS3 yield of 41.3% was obtained when annealing was carried out at 400 MPa/60°C for 15 min, whereas it took nearly 8 h to obtain the same yield under conventional atmospheric annealing at 60°C. The yield of RS3 could be further significantly increased by annealing under 400MPa/60°C pressure for 15 min followed by resting at atmospheric pressure for 3 h 45 min, and repeating this cycle for up to six times. Microstructural surface analysis of the product under a scanning electron microscope showed an increasingly rigid density of the crystalline structure formed, confirming higher RS3 content.

Keywords: Debranched cassava starch; High pressure annealing treatment; Type 3 resistant starch

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1. Introduction

Resistant starch (RS) is the non-digestible starch, which can resist the digestion by $\alpha$-amylase and act like dietary fibres - that helps to promote the growth of beneficial bacteria in the intestine (Englyst, Kingman, & Cummings, 1992). Depending on its botanical origin and process employed to form it, RS can be divided into four categories: RS1 is physically inaccessible for reasons such as starch entrapment in a protein matrix or a plant cell wall (e.g. in seeds and unprocessed whole grain); RS2 is raw granular starch which cannot be absorbed by small intestine (e.g. those from potato and green banana); RS3 is retrograded starch, mainly retrograded amylose formed during cooking and cooling processes; and RS4 is chemically modified starch which is cross-linked by chemical agents and insusceptible to digest and absorb in the small intestine (Chung, Donner, & Liu, 2011).

In general, many methods involving physical, chemical and enzymatic transformations have been employed to alter the properties of starch, which enhance health attributes and/or minimize defects in structure. Researchers have attempted to improve the RS yields by: 1) heat-moisture treatment and annealing (Brumovsky & Thompson, 2001), 2) enzyme treatment (Vatanausuchart, Tungtrakul, Wongkrajang, & Naivkul, 2010; H. Zhang & Jin, 2011), 3) combined heat/enzyme treatment (Mutungi, Rost, Onyango, Jaros, & Rohm, 2009) and 4) chemical treatment (Haynes et al., 2000). Since consumers are increasingly interested in natural and organic foods for health and environmental reasons, employing physical and/ or enzymatic treatment appears more attractive.

Although gelatinized starch retrogrades upon cooling and typically has a negative effect on the quality of starchy foods, low-temperature storage leads to the formation of retrograded starch or RS3 fraction. RS3 is of particular interest as a food ingredient because of its physical and nutritional functionality and processing stability (Thompson, 2000). Several factors influence
the quality and quantity of RS3 in addition to storage conditions: amylose (a substantially linear glucose polymer) and amylopectin (a mainly branched glucose polymer) ratio, length of polymer chains or degree of polymerization (DP), retrogradation or recrystallization of amylose, water content, processing steps/conditions, and the presence of lipid and other components influencing gelatinization and/or the retrogradation process (Eerlingen & Delcour, 1995). In addition, there are a number of studies have demonstrated that the higher the level of amylose, the greater the RS3 fraction formed (Brown, Mcnaught, & Moloney, 1995). As a consequence, using a debranching enzyme like pullulanase to act on gelatinized starch has become one of the most important methods employed to directly cleave the branches of amylopectin and produce linear α-glucan chains (Pongjanta, Utapattanaceep, Naivikul, & Piyachomkwan, 2009; Vatanasuchart et al., 2010; H. Zhang & Jin, 2011).

Based on studies involving physically modified starches, research has indicated that the high hydrostatic pressure (HHP) plays an important role in inducing gelatinization of starches while still maintaining their granular integrity (Kasemwong, Ruktanonchai, Srinuanchai, Itthisoponkul, & Siroth, 2011; Oh, Pinder, Hemar, Anema, & Wong, 2008). However, subsequent to HHP treatment, a rapid retrogradation has been observed (Kawai, Fukami, & Yamamoto, 2007). Additionally, a combination of pressure and temperature can create more nuclei in starches (Hartel, 2001) and lead to a higher yield of recrystallized starch as a resistant starch product. This treatment was used for wheat starch to produce resistant starch following several steps such as annealing and storage, enzyme/acid hydrolysis and annealing-pressure cycle (Bauer, Wiehle, & Knorr, 2005). It has been reported that a combination of treatments enhances the yields of RS in wheat starch more than by using individual processes such as HHP or thermal treatment.

Native cassava or tapioca starch (Manihot esculenta Crantz) is one of the food ingredients consisting of 17% amylose and 83% amylopectin (Breuninger, Piyachomkwan, & Siroth, 2009). Given its high degree of branching, a higher formation of RS3 fraction may be expected by
debranching (Mutungi, Onyango, Jaros, Henle, & Rohm, 2009; Vatanasuchart et al., 2010).

Furthermore, there are many methods available to improve the recrystallization of debranched cassava starch, such as by annealing, autoclaving-cooling cycle and/or heat-moisture treatment resulting in the rise of RS3 content (Mutungi, Rost, et al., 2009).

Despite its promising potential, very limited information exists within published scientific literature on the effects of high hydrostatic pressure (HHP) on retrograded resistant starch or the RS3 content of debranched starch, particularly from cassava or tapioca. This study explores the potential of forming RS3 in the structure resulting from the recrystallized debranched cassava starch, following the application of combined HHP and thermal annealing. This work also aims to evaluate the use of HHP within a solvent-free environment to maximise the production of RS3.

2. Materials and methods

2.1. Materials

Native cassava starch was supplied by Siam Modified Starch Co., Ltd (Thailand) in the form of white powder, containing less than 14% moisture and 0.2% ash content. Pullulanase solution (Sigma E2412, 1,824.68 U/ml), Trehalose (Fluka 90208), Maltotetraose (DP4 - Supelec 47877), Maltopentaose (DP5 – Supelec 47876), Maltohexaose (DP6 – Supelec 47873), Maltoheptaose (DP7 – Supelec 47872) were all obtained from Sigma-Aldrich Co., Ltd. (United Kingdom). Resistant starch assay kit (K-RSTAR) was purchased from Megazyme International Ireland Ltd. (Ireland). All other chemicals was used were of analytical grade.

2.2. Process steps to produce resistant starch from cassava starch
2.2.1. Debranching of cassava starch

Native cassava starch (NS) was hydrolysed by pullulanase enzyme to cleave the α 1,6 glycosidic bonds at the branched points of amylopectin molecules; the method employed was adapted from Mutungi, Rost, et al. (2009). A stock solution of pullulanase (Sigma E2412) was diluted in 20 mM sodium acetate-hydrochloric acid buffer (pH 5.0) to 25 U/ml before use. The starch (20 g) was weighed into 250 ml polycarbonate centrifuge bottle and suspended in 140 ml of 20 mM sodium acetate-hydrochloric acid buffer (pH 5.0). The starch suspension (12.5%w/w) was gelatinized by autoclaving at 121°C for 15 min and cooled to 50°C. The starch gel was mixed with 20 ml of 25 U/ml pullulanase solution. This mixture (11%w/w starch) was incubated at 50°C in a shaking water bath (Grant OLS 200, Cambridge, UK) at 100 stroke/min in linear motion for 24 h. The residue was recovered after inactivating the enzyme by washing thrice with chilled deionized water (temperature < 4°C) and centrifuging for 10 mins at 3000 rpm (Sorvall RC-5B Plus, Kendro, Newtown, USA). The use of chilled water for inactivating this enzyme has been employed earlier by Mutungi, Rost, et al. (2009). The pellet was freeze-dried (Martin Christ Gamma 2-16, Osterode am Harz, Germany), ground in a mortar, and sieved through a mesh size of 212 µm. This sample was termed DS.

2.2.2. Incubation of debranched-autoclaved cassava starch

The effect of incubation condition on the yields of RS3 from the debranched-autoclaved cassava starch (DAS) was determined. A 2 x 5 x 5 factorial experiment covering starch solution concentration, temperature and time, was performed and carried out in triplicates. The DS (0.5 g) was weighed into 30 ml glass vial and mixed with 4.5 and 2 ml deionized water to form 10 and 20%(w/w) concentrated solutions, respectively. The mixtures were autoclaved at 121°C for 15 min and cooled to 50°C. These samples were allowed to stand for 15 min at ambient temperature before incubating at 4°C (in a fridge), 20, 50, 60 and 90°C (in water baths) for 0.25,
2, 4, 8, and 24 h. The two sets of solutions (i.e. 10 and 20%) were freeze-dried (VirTis Bench Top K Series, SP Industries, Warminster, PA, USA). These samples were referred to as: **DAS-10** (i.e. the one from 10% DS) and **DAS-20** (from the 20% DS).

### 2.2.3. Pressurizing of debranched-autoclaved cassava starch

The effect of high hydrostatic pressure annealing on the development of RS3 content was investigated. The debranched starch (DS, 0.5 g) was weighed into 30 ml glass vial and mixed with 2 ml deionized water and autoclaved at 121°C for 15 min to make the DAS-20. The resulting gel was transferred to a polyethylene pouch; vacuum packed (Mutivac A300, Wolfertschwenden, Germany), and pressurized in a high-pressure vessel (37mm diameter and 246 mm length) (Stansted Fluid Power type Food Lab 900, Stansted, U.K), where the temperature was controlled at 60°C by using a circulating thermostatic water bath (Grant B20-632, Cambridge, UK). Two sets of experiments were undertaken in the high pressure rig:

In the first set of experiments, the pressure applied was constant and continuous for a given period of time. Packed samples of DAS-20 were pressurized at 200, 400 and 600 MPa for 0.25, 0.5, 1, 2, 4, 8 and 24 h at 60°C to yield a 3 x 7 factorial experiments, each performed in triplicates. The samples were then unpacked and transferred to 15 ml centrifuge tube before being freeze-dried. These samples were named **HPT-DAS**.

In the second set of experiments, the application of high pressure was intermittent or cyclic. The DAS-20 samples were subjected to pressure of 400 MPa at 60°C for 15 min followed by atmospheric holding for 3 h and 45 min constituting one cycle; this cycle was repeated up to six times covering a total treatment period of 24 h. Samples were drawn for analysis after each cycle, and RS3 contents were compared with a corresponding control sample, which was simply incubated at atmospheric pressure and 60°C for the same duration of time. This set of experiments therefore involved a 2 x 6 factorial performed in triplicates. The cyclic treated
samples, named **HPC-DAS**, were then unpacked, transferred to 15 ml centrifuge tube and freeze-dried, individually.

2.3. *Measurement of chain length distribution of debranched starch*

High-performance anion exchange chromatography, equipped with pulsed amperometric detector (HPAEC-PAD) and a CarboPac PA200 Dionex DX-600 (Dionex, Sunnyvale, CA, USA) was undertaken to determine polymer chain length distribution of the debranched cassava starch. The sample prepared and condition employed were adapted from Dionex corporation (2004) and Mutungi, Rost, et al. (2009). Trehalose (10 mg) was suspended in 10 ml of ultrapure water (UPW) and used as an internal standard. Molto-oligosaccharide standards (DP4 – DP7, 1 mg/ml) were prepared in 150 mM aqueous sodium hydroxide solution and instantly diluted 50-fold with UPW, containing 10 µl of internal standard stock. The standards were performed to identify peak by comparing retention time of sample peaks with those of standards, and to predict peaks at the higher DP7 of samples according to the linear relationship between the retention time and the degree of polymerization. Debranched starch sample (DAS-20, 20 mg) was weighed into 2 ml vials to which 400 µl of 2 M aqueous sodium hydroxide was added and mixed in a vortex mixer. This suspension was then diluted with 1600 µl of ultrapure water and mixed in the same vortex mixer at 4°C and 450 rpm for a further period of 24 h. A 20 µl aliquot of the solution was diluted 50-fold with 980 µl 150 mM aqueous sodium hydroxide solution/10 µl internal standard stock. All samples were filtered through a 0.2 µm filter and 25 µl was autoinjected at 0.5 ml/min flow rate into the column. The waveform and durations applied were as follows: $E_1 = 0.1V \ (t_1 \ 0 \ s)$, $E_2 = 0.1V \ (t_2 \ 0.20 \ s)$, $E_3 = 0.1V \ (t_3 \ 0.40 \ s)$ (integration from 0.2 to 0.40 s), $E_4 = -2.0V \ (t_4 \ 0.41s)$, $E_5 = -2.0V \ (t_5 \ 0.42 \ s)$, $E_6 = 0.6V \ (t_6 \ 0.43 \ s)$, $E_7 = -0.1V \ (t_7 \ 0.44 \ s)$ and $E_8 = -0.1V \ (t_8 \ 0.50 \ s)$. A gradient of 100 mM sodium hydroxide solution (Eluent A) and 150 mM sodium hydroxide solution, containing 500 mM sodium acetate (Eluent B) was used for
elution. Increasing concentration of eluent B from 5-40% (0-13 min), 40-85% (13-50 min) and
decreasing to 5% (50-70 min) were applied in linear gradients. Integrating area under individual
peaks was determined by using Chromleon® version 6.6 software (Dionex). This experiment
was performed in triplicates.

2.4. Determination of resistant starch

The amount of resistant starch (RS) in all samples was investigated in triplicates using resistant
starch assay kit (Megazyme, Bray, Ireland) - an enzymatic method recommended by the
Association of Official Analytical Chemists (AOAC) Method 2002.02 (McCleary & Monaghan,
2002). The main features of this procedure are: removal of non-resistant starch by hydrolysis
and solubilisation using pancreatic α-amylase and amyloglucosidase (AMG), washing the
residue with ethanol, neutralization and enzymatic hydrolysis of RS to glucose using 2M KOH
acetate buffer and AMG, and measurement of RS by quantification of glucose with glucose
oxidase/peroxidase reagent (GOPOD). The RS was calculated as mg glucose × 0.9.

2.5. Evaluation of microstructure

The surface of the resistant starch samples were scanned using scanning electron microscope or
SEM (S360, Leica Cambridge, UK). A small amount of dried sample was attached to
electrically conductive double-sided adhesive carbon disc, which was pressed on a specimen
stub. Gold was used to coat the sample using a sputter coater (S150B, BOC Edwards, Crawley,
UK). The SEM operation conditions were: working pressure < 1.0E-4 Torr, accelerating
voltage = 20 kV and working distance = 14 mm at the magnifications of 300×, 1000× and
3000\times \). The image was recorded using IScan 2000 image software (ISS Group, Manchester, UK).

2.6. Statistical analysis

All RS percentages obtained were subjected to analysis of variance – ANOVA using PASW statistics 18.0 software (SPSS, IBM, Somer NY, USA). The results were expressed as mean values with standard deviation. The differences between the group mean values were established at 95% confidence interval \((P < 0.05)\) using Duncan’s new multiple range test (DMRT).

3. Results and Discussion

3.1. Chain length distribution of debranched starch

Native cassava starch was debranched using pullulanase and the chain length distribution of debranched starch was measured by high performance anion exchange chromatography equipped with pulsed amperometric detector (HPAEC-PAD). According to Hanashiro et al. (1996), branch chain types of amylopectin are classified by HPAEC to the group with periodicity of 12 as DP 6–12, 13–24, 25–36 and DP ≥ 37. These ranges of DP are referred to as A-chains, B\(_1\)-chains, B\(_2\)-chains and B\(_n(\geq 3)\)-chains, respectively. In the present study, a polymer chain distribution between DP 4-45 was obtained and it is illustrated in Fig. 1. Most of the shorter chains were removed with cold water, showing the amount of chains of DP ≤ 5 remaining, to be only 0.5 ± 0.2 %. The proportion of A-chains (DP 6-12) was 23.0 ± 0.6 %. The highest yield of B\(_1\)-chains (DP 13-24) was found to be 50.6 ± 1.1 %, whereas the B\(_2\)-chains (DP 25-36) and B\(_3\) to B\(_4\)-chains (DP ≥ 37) were lower at 22.0 ± 0.3 % and 3.9 ± 0.6 %, respectively.

These results indicate that the cassava amylopectin mainly comprises of A and B\(_1\)-chains, which
conforms to the literature results, even though it varies with the cultivars (Charoenkul, Uttapap, Pathipanawat, & Takeda, 2006; Mutungi, Rost, et al., 2009). The average chain length is DP 20.5, which is similar to the values observed in previous studies for debranched cassava starch (Mutungi, Rost, et al., 2009; Tester, Karkalas, & Qi, 2004). Schmiedl et al. (2000) also reported that effective formation of RS3 can be produced from linear glucose chains of DP 10-35.

3.2. Effect of debranching and autoclaving on the formation of RS3

The RS contents of native starch (NS), debranched starch (DS), and debranched-autoclaved starch (DAS) are presented in Table 1. After debranching, the amount of RS increases drastically from 2.4 ± 0.2% in NS to 17.4 ± 0.5% in DS. These results are also consistent with other studies that demonstrated debranching by using pullulanase enzyme in: 1) maize starch, which increased the RS yield from 0.60 to 25.5% within 24 h (Marija, Milica, & Ljubica, 2010); and 2) corn starch, where RS yield increased from 0.67 to 19.02% within 12 h (Gao, Li, Jian, & Liang, 2011). This implies that the hydrolysis of α-1 →6 linkages in amylopectin can produce more linear structures similar to the amylose chains, and/or create free A-chains of amylopectin in the form of double helix and crystallite segments. These debranched structures closely pack into the crystal formation as retrograded starch (RS3) during retrogradation or the annealing period (Vasanthan & Bhattiy, 1998).

Moreover, the results given in Table 1 confirm that the RS content of DS is higher than NS. It may be mentioned here that there are conflicting reports in literature about the RS contents of DS and NS. Mutungi, Rost et al. (2009) found that the RS content in DS (21.43g/100g) was significantly lower compared to that in NS (43.96g/100g), which was also observed by Vatanasuchart et al. (2010). In contrast, Charles et al. (2005) noted that the initial RS yield in five native cassava starches was only 6.8–14 %. These discrepancies may be due to the different
botanical origin of cassava. It may also be noted that the RS fraction in NS is type 2 due to the compact structure limiting the accessibility of the digestive enzyme in granular starch form, whereas in DS, it is type 3 which results in retrograded polymer chains being formed in the gelatinized starch (Mutungi, Rost, et al., 2009; Ozturk, Koksel, Kahraman, & Ng, 2009).

After autoclaving of 10 and 20 %w/w DS, the RS3 content of debranched-autoclaved starch (DAS-10 and DAS-20) is significantly higher (22.0 ± 0.5 and 28.3 ± 1.0%, respectively) than that of DS (17.4 ± 0.2%). During the autoclaving experiment, the temperature was gradually increased to 121°C, maintained steady for 15 min, and then cooled down to 50°C before taking samples. The total time for this process was approximately 2 h. Thus, the RS3 forms in two steps that involve starch hydrolysis during autoclaving at a high temperature, followed by recrystallization during cooling. It is important to note that the proportion of RS3 is strongly enhanced when starch is debranched to increase the number of linear molecules prior to thermal treatment (i.e. autoclaving). A similar response was also seen in the case of debranched-autoclaved wheat starch (Berry, 1986) and banana starch (González-Soto, Agama-Acevedo, Solorza-Feria, Rendón-Villalobos, & Bello-Pérez, 2004).

3.3. Effect of concentration, temperature and time on the formation of RS3

Fig. 2 shows the proportion of RS3 in debranched-autoclaved cassava starch of two concentrations (10 and 20%w/w of DS) at various temperatures (4, 20, 50, 60 and 90°C), and times (0, 0.25, 2, 4, 8 and 24 h). The results suggest that there are significant interaction effects ($P < 0.05$) between concentration, temperature and treatment time. The initial RS3 content of debranched-autoclaved starch at the higher concentration DAS-20 (28.3 ± 1.0% RS) is clearly higher than DAS-10 (22.0 ± 0.5% RS). Thus, DAS-20 is more effective in recrystallization due to re-association of a significant amount of short linear α-glucan occurring with deionized water
as the plasticizer. In contrast, DAS-10 has excessive water, leading to an obstruction of the intermolecular interaction between the hydrogen bonding of short chain fragments (Y. Zhang & Rempel, 2012).

After incubating, the RS3 contents in DAS-10 and DAS-20 are found to follow a similar trend. The yield of RS gradually rises from 4 to 60°C, and then drops at 90°C. Theoretically, the mechanism of recrystallization in amorphous polymers consists of nucleation, propagation, and maturation – which represents crystal perfection by slow growth. The nucleation rate of linear glucans largely increases as the temperature decreases to the glass transition temperature \( T_g \) at approximately -5°C, whereas the propagation rate increases as the temperature increases to the melting temperature \( T_m \) of about 150°C (Biliaderis, 2009; Ring et al., 1987). In other words, the recrystallization correlates with the molecular mobility and crystal growth rate, and can occur between \( T_g \) and \( T_m \) or in the glassy state (Marsh & Blanshard, 1988). Consequently, the treatment temperature at 4 and 20°C may be above and close to the \( T_g \), which favours nuclei formation, however, the crystals tend to develop slowly. Thus, the increased RS3 yield of DAS-10 and DAS-20 at low temperature is not observed even after a prolonged incubation time to 24 h. The RS formation at 90°C also reveals a distinctively low value within 24 h at both concentrations; this indicates that the temperature may be above \( T_m \), which means the glucan polymers are completely transferred into the liquid state. Therefore, there is no positional order in the short chain molecules to result in the formation of RS. On the other hand, the formation of increased RS3 in the case of DAS-10 and DAS-20 at 50 and 60°C, further increased with treatment time. The highest RS3 content of 29.6 ± 0.5 % in DAS-10 and 36.5 ± 0.3 % in DAS-20 is accomplished at 60°C after 4 h. Clearly this temperature influences orientational mobility and formation of double helices to promote crystal growth for retrograded starch (Jayakody & Hoover, 2008). In summary, based on the present study, the highest RS3 content is formed by the autoclaving of 20% debranched starch solution at 60°C for 4 h.
3.4. Effect of high-pressure processing and annealing on the formation of RS3

Fig. 3 shows the effect of high-pressure treatments at 60°C on the yield of RS3 in DAS-20. It is clear that the RS3 levels in DAS-20 can significantly improve by combining pressurizing at 400 MPa and annealing for various times (Fig. 3A); although at a treatment time of 24 h, the increase in RS3 is not significantly different \((P > 0.05)\) between 400 MPa HPT-DAS and the control-annealed sample. During the first 15 min, the RS3 formation in the combined treatment at 400 MPa surprisingly accelerated with a 25.7% increase (from 28.7 ± 0.5% to 36.1 ± 0.5% RS), whereas the control-annealed sample only had a 9.7% increase of RS (from 28.7 ± 0.5% to 31.5 ± 1.1% RS). These results clearly indicate that pressuring for short times provides a significantly higher initial rate of recrystallization.

To obtain a higher yield of RS3, DAS-20 samples were subjected to cyclic high pressure annealing treatment. As the results demonstrate, cyclic pressure annealing at prolonged treatment times (Fig. 3B) can further enhance the RS3 formation. Although during the first 12 h the increase in RS3 was not significantly different \((P > 0.05)\) between HPC-DAS and HPT-DAS at 400 MPa, after four to six cycles (16-24 h) the RS3 yield of HPC-DAS was greater than that of either the 400 MPa HPT-DAS and the control-annealed samples. A possible explanation is that, the seed crystal formation occurs when the volume of the system decreases by increasing pressure, forcing starch molecules closer together and creating more nuclei in the glassy state. Although high-pressure application accelerates the nucleation rate, these nuclei are limited to propagate. However, these seeds lead to the formation of large crystals when holding at atmospheric pressure (lower pressure level). This scenario is similar to the high-pressure crystallization of cumin aldehyde (essential oil) \((\text{Moritoki, Nishiguchi, \\& Nishida, 1997})\) and lysozyme (protein) \((\text{Moritoki, Nishiguchi, \\& Nishida, 1995})\). Therefore, it implies that the
propagation of crystalline DAS is restricted under high-pressure but cyclic pressure annealing improves the rate of propagation.

On the other hand, despite an increase in RS, HPT-DAS at 200 and 600 MPa were not significantly different in RS3 content between the groups, and their RS contents were lower than the control sample even after pressurizing for 24 h (Fig. 3A). This behaviour is probably due to limiting propagation of crystalline DAS under high-pressure conditions at 200 MPa, while the low level of RS3 at 600 MPa could be due to the crystalline melting of DAS. A number of published papers report partial melting of the crystalline structure during compression at very high pressure: for instance, at 650 MPa in polylactides (Ahmed, Varshney, Zhang, & Ramaswamy, 2009); and at 740 – 1,500 MPa in several native starches, including normal corn, waxy corn, wheat and potato starches (Liu, Selomulyo, & Zhou, 2008).

Overall, the high hydrostatic pressure conditions for producing highest amount of RS3 are suggested to be concurrently pressurizing DAS-20 at 400 MPa and annealing at 60°C for 15 min. Clearly these conditions significantly reduce the process time, from 8 h (single incubating at 60°C) to only 15 min with the same RS content generated (36% RS) as in the case of the conventional process. The highest RS yield was obtained after six cycles (24 h of total time) of pressure alternating between 400 MPa and atmosphere under the temperature of 60°C (41.9 ± 0.5% RS). Bauer et al. (2005) also noted increase in RS content from 2 to 12 % in the case of wheat starch when the pressure was increased to 500 MPa for 15 min every 24 h, over a period of 10 days. Thus, it is important to note that high-pressure application on debranched starch results in greater enhancement of RS compared to high-pressure treated native starch that still maintains its granular form.

3.5. Microstructure of RS3
The effects of: debranching and autoclaving on cassava starch (Fig. 4), concentration, temperature and time (Fig. 5A), and high-pressure treatments on autoclaved samples (Fig. 5B) were monitored using scanning electron microscopy (SEM). Fig. 4A shows the native starch (NS) in a granular form. After debranching, the NS loses its granular structure, and then the surface of DS appears more fluffy which indicates that the glucan polymers only reassociate loosely (Fig. 4B). After autoclaving the DS, a densely packed surface region is evident in DAS (Fig. 4C). In Fig. 5A, it is apparent that the DAS-20 incubation at 60°C for 4 h exhibits a smoother area than DAS-10. When subjecting DAS-20 to high pressure annealing treatments, Fig. 5B demonstrates that HPC-DAS after six cycles (24 h of process time) has a more densely packed surface than HPT-DAS. It clearly shows less porosity and a smoother surface area, which would yield greater resistance to enzyme digestion and increasing the RS content. The overall microstructural observations show that RS3 content increases as the rigid dense crystalline structure increases.

4. Conclusion

The process employed plays an important role in accelerating retrogradation and the transforming of native starch into RS3. In this study, the debranching step gave more linear glucans and the autoclaving step aggregated these to increase crystallinity. High pressure annealing subsequently accelerated RS formation within 15 min, in contrast to atmospheric annealing (single incubation) which required up to 8 h to result in the same yield of RS3. Thus, process times can be drastically reduced using high pressure annealing. Yields of RS3 fraction can be further increased following cyclic high pressure annealing of debranched-autoclaved starch. The highest RS yield was obtained after applying six cycles (24 h of process time) of pressure, each alternating between pressure application (400MPa/60°C/15 mins) to accelerate
the nucleation rate of starch crystallization, and incubation (atmospheric pressure/60°C/3h 45 mins) for crystal propagation (41.9 % RS). These conditions gave the highest yield of RS3 from a 20% w/w solution of debranched-autoclaved starch. Thus, the high pressure annealing treatment is highly promising to increase RS yield. In addition, this method intensifies the formation of RS by physical modification (i.e. without using solvents), which is safer for food industry use.

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Reference


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**Fig. 1** Polymer chain length distribution of debranched amylopectin of cassava starch (DS) using high performance anion exchange chromatography with pulsed amperometric detector (HPAEC-PAD). The DP4-45 is shown in the chromatogram. DP4-7 peak labels indicate the DP from molto-oligosaccharide standards. The inset shows the relative peak area for the individual DP from the mean of three independent measurements.
Fig. 2 Effect of concentration (10 and 20% w/w of debranched starch), temperature (4, 20, 50, 60 and 90°C) and time (0, 0.25, 2, 4, 8 and 24 h) on yield of RS3 (% w/w) of debranched-autoclaved cassava starch (DAS).
Fig. 3 Effect of high-pressure incubation on yield of RS3 from debranched-autoclaved starch (DAS-20, 20% w/w of debranched starch) (A) high pressure (200, 400 and 600 MPa) at 60°C for different times (0.25, 0.5, 1, 2, 4, 8 and 24 h): HPT-DAS. The inset to A shows increasing RS3 content on an expanded scale. (B) high pressure incubation at 400 MPa and 60°C for 15 min followed by atmospheric holding at the same temperature for 3h 45 min, repeating this cycle for up to six times: HPC-DAS. Figure B also shows RS content comparison among control treatment (atmospheric annealing at 60°C), 400 MPa HPT-DAS and 400 MPa HPC-DAS, at any treatment time.
**Fig. 4** Scanning electron micrographs at magnification of 300 × and 3000 × of (A) native cassava starch: NS, (B) debranched starch: DS, and (C) debranched-autoclaved starch at 20%w/w of DS: DAS-20
Fig. 5 (A) Scanning electron micrographs at magnification of 300 × and 1000 × after incubation at 60°C for 4 h of (A1) debranched-autoclaved starch at 10%w/w of DS: DAS-10, and (A2) debranched-autoclaved starch at 20%w/w of DS: DAS-20. (B) Scanning electron micrographs at magnification of 300 × and 1000 × of debranched-autoclaved starch at 20%w/w of debranched starch after applying high pressure (B1) high pressure annealing treatment at 400 MPa and 60°C at a treatment time of 24 h: HPT-DAS, and (B2) cyclic high pressure annealing at 60°C after six cycles with pressure swinging between 400 MPa for 15 min and atmosphere for 3 h 45 min - over 24 h: HPC-DAS.
Table 1: Effect of debranching and autoclaving on RS3 content of cassava starch

<table>
<thead>
<tr>
<th>Samples</th>
<th>Resistant starch (%)</th>
<th>Starch sources</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>NS</td>
<td>2.4 ± 0.2&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Siam Modified Starch Co. Ltd, Thailand</td>
<td></td>
</tr>
<tr>
<td>DS</td>
<td>17.4 ± 0.5&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DAS-10</td>
<td>22.0 ± 0.5&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DAS-20</td>
<td>28.3 ± 1.0&lt;sup&gt;d&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NS</td>
<td>43.9</td>
<td>Kenya Industrial Research and Development Institute, Kenya</td>
<td>Mutungi, Rost, et al., 2009</td>
</tr>
<tr>
<td>DS</td>
<td>21.4 ±2.7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NS</td>
<td>58.2 ± 1.3</td>
<td>Taiwa Public Co., Ltd, Thailand</td>
<td>Vasanthan &amp; Bhatti, 1998</td>
</tr>
<tr>
<td>DS</td>
<td>13.0 ± 1.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NS</td>
<td>6.8 – 14.0</td>
<td>5 cassava genotypes (Rayong2, Rayong5, KU50, Hanatee and YOO2), Thailand</td>
<td>Charles, Chang, Ko, Siroth, &amp; Huang, 2005</td>
</tr>
</tbody>
</table>

<sup>a-d</sup> Mean ± standard deviation followed by the different superscripts are significantly different (P < 0.05). NS: Native starch; DS: Debranched starch; DAS-10: Debranched-autoclaved starch at 10%DS; and DAS-20: Debranched-autoclaved starch at 20%DS