**Xanthosoma riedelianum starch for use in the food industry**

**Abstract** – The objective of this work was to characterize the starch of *Xanthosoma riedelianum*, commonly known as “mangarito” in Brazil, and to evaluate its properties for the food industry. The starch was characterized as to its morphology and its thermal and technological properties. Morphology was evaluated by the analysis of granule size distribution, scanning electron microscopy, and X-ray diffraction. The thermal properties were examined by determining the initial, peak, and conclusion gelatinization temperatures, whereas the technological properties were obtained by analyzing viscosity, swelling power, solubility index, paste clarity, and syneresis. Although the extraction of *X. riedelianum* starch showed a low yield, the species is a source of starch with a high sticking temperature, mechanical stirring resistance, tendency to retrograde, and final viscosity. Therefore, this starch is ideal for products that require high viscosity, such as pie fillings, puddings, children’s foods, and bakery products, and its paste, which is opaque and viscous, can also be used in the formulation of broths.

**Index terms**: rheological properties, thermal properties, unconventional vegetable.

**Amido de Xanthosoma riedelianum para uso na indústria de alimentos**

**Resumo** – O objetivo deste trabalho foi caracterizar o amido de *Xanthosoma riedelianum*, comumente conhecido como mangarito no Brasil, e avaliar suas propriedades para a indústria alimentícia. O amido foi caracterizado quanto à sua morfologia e às suas propriedades térmicas e tecnológicas. A morfologia foi avaliada por meio de análises de distribuição de tamanho dos grânulos, microscopia eletrônica de varredura e difração de raios X. As propriedades térmicas foram examinadas por meio da determinação das temperaturas inicial, de pico e de conclusão de gelatinização, enquanto as propriedades tecnológicas foram obtidas pelas análises de viscosidade, poder de inchamento, solubilidade, claridade da pasta e sinérese. Embora a extração do amido de *X. riedelianum* tenha apresentado baixo rendimento, a espécie é uma fonte de amido com altas temperaturas de empastamento, resistência à agitação mecânica e tendência à retrogradação, bem como elevada viscosidade final. Portanto, o amido é indicado para produtos que requerem elevada viscosidade, como recheios de tortas, pudins, alimentos infantis e produtos de panificação, e sua pasta, que é opaca e viscosa, também pode ser usada na formulação de caldos.

**Termos para indexação**: propriedades reológicas, propriedades térmicas, hortaliça não convencional.
Introduction

Starches have been modified by the food industry through costly processes – chemical, physical, or enzymatic, for example – that alter their typical properties (Lewicka et al., 2015). However, the food sector has been increasing the search for native starches with specific properties, such as a greater resistance to light and strong mechanical stresses, high acidity, and compatibility with the ingredients present during the cooking phase, including acids, sugars, fats, and salts (Waterschoot et al., 2015).

*Xanthosoma riedelianum* (Schott) Schott, commonly known in Brazil as “mangarito”, is an example of a species that stands out nutritionally as a highly energetic source of starch, with 95.19 g 100 g⁻¹ starch stored in its rhizomes (Ávila et al., 2012). The starch from this species has a caloric index of approximately 100 kcal 100 g⁻¹, a dry matter content between 17 and 20%, and a protein content from 3.0 to 3.5% (Ávila et al., 2012; Madeira et al., 2015). Currently, the market for *X. riedelianum* is on the rise due to the growing interest in haute cuisine and to the great potential of the species for family farming, attributed to its rusticity, low production cost, and profitability (Madeira et al., 2015).

Therefore, starch is relevant for Brazilian agribusiness, especially for family farming, showing the importance of the rescue of unconventional species by the market and, consequently, of studies on their production, considering the limited technical information available in the literature about their cultivation (Souza et al., 2018). Incipient studies on *X. riedelianum* should focus on its increased productivity and commercial quality (Madeira et al., 2015).

Research on the technological, physicochemical, and morphological properties of starches from unconventional sources is essential to improve the functionality of these substances and to verify their applicability in the food, pharmaceutical, textile, packaging, and paper industries (Souza et al., 2019; Macena et al., 2020).

The objective of this work was to characterize the starch of *Xanthosoma riedelianum*, commonly known as “mangarito” in Brazil, and to evaluate its properties for the food industry.

Materials and Methods

The experiment was conducted at the Department of Food Science of Universidade Federal de Lavras, located in the municipality of Lavras, in the state of Minas Gerais, Brazil. The *X. riedelianum* rhizomes used for starch extraction were supplied by Embrapa Hortaliças (Brasilia, DF, Brazil). The experimental design was completely randomized, with three replicates for each analysis.

The starch extraction process followed the recommendations of Daiuto & Cereda (2003). The material was oven-dried at 45°C for 24 hours. The starch extraction yield (EY), expressed in percentage, was calculated by the equation:

\[
\text{EY} = \frac{\text{Starch weight (g)}}{\text{Weight of } X. \text{ riedelianum with peel}} \times 100
\]

Humidity was determined by the IV3000 infrared light meter (Gehaka, São Paulo, SP, Brazil).

Common-light and polarized images of the starch granules were obtained by the BX51-p optical microscope (Olympus, Tokyo, Japan). The particle size distribution of the starch was directly determined by laser diffraction spectroscopy on the Mastersizer 3000 particle size analyzer (Malvern Panalytical Ltd, Worcestershire, United Kingdom). The average diameter of the granules was measured using the diameter of a sphere of the same volume and the De Brouckere mean diameter, also know as volume-weighted mean diameter \[D (4.3)\] (Jinapong et al., 2008). The size of the starch granules was determined through the LEO EVO 40 XVP scanning electron microscope (Leo Electron Microscopy Ltd, Cambridge, United Kingdom) at 20 kv and a distance between 7 and 15 mm (Fernandes et al., 2019).

In order to analyze starch organization, the PANalytical X’Pert PRO X-ray diffractometer (Malvern Panalytical Ltd, Worcestershire, United Kingdom) was used, with Co radiation and Fe filter according to Hayakawa et al. (1997). Starch crystallinity was quantitatively determined, and the curve connecting the peak bases was plotted on the diffractograms, in addition to a linear base. The area between the curve and the linear base of the diffractogram corresponds to the amorphous area. Relative crystallinity was obtained by the ratio between the crystalline area and the total area shown in the diffractograms, being presented in percentage. The degree of relative crystallinity was estimated using the Microcal Origin, version 8.6, software (OriginLab Corporation, Northampton, MA,
USA), following the method described by Barbi et al. (2018).

To determine thermal properties, the differential scanning calorimetry (DSC) analysis was performed using the DSC-60A equipment (Shimadzu Corporation, Kyoto, Japan). For this, 4 mg of starch at a dry basis were added to 6 μL deionized water, and the samples were kept for 2 hours at room temperature for balance and then placed in a calorimeter and heated to 5°C per minute, from 25 to 100°C (Leonel et al., 2011). Initial, peak, and conclusion temperatures, as well as the enthalpy variation of the starches, were determined using the TA-60WS thermal analysis workstation (Shimadzu Corporation, Kyoto, Japan). Thermogravimetry (TGA) was measured on the DTG-60A/60AH equipment (Shimadzu Corporation, Kyoto, Japan), and the TA-60WS thermal analysis workstation (Shimadzu Corporation, Kyoto, Japan) was used to obtain the values observed on the curves. Viscosity drop (breakdown) and tendency to retrograde (setback) were also assessed.

Paste properties were analyzed with the rapid viscosity analyzer Rapid Visco Analyzer (Newport Scientific Pty Ltd, Warriewood, Australia), in order to determine the viscosity profile of the starches. The values for paste temperature and for maximum, minimum, and final viscosity, as well as for viscosity drop and tendency to retrograde, were obtained according to Silva et al. (2013).

Swelling power and the solubility index were determined by the methodology adapted from Leach et al. (1959), at room temperature and at 60, 70, 80, and 90°C. Paste clarity was evaluated following the protocol described by Craig (1989) using 1 g 100 mL⁻¹ starch suspensions that were gelatinized for 30 min in a boiling water bath. Transmittance was determined at 650 nm with the Varian Cary 50 spectrophotometer (Agilent Technologies, Inc., Santa Clara, CA, USA). The samples were stored at 4°C for eight days in order to monitor retrogradation, and transmittance was assessed every 24 hours from the first to the eighth day.

Syneresis was obtained by the method proposed by Singh et al. (2004), being considered the amount of water released (%), according to the equation:

\[
\text{Syneresis (\%)} = \frac{\text{Released liquid (g)}}{\text{Weight of the initial sample (g)}} \times 100
\]

Means and standard deviations were calculated for the results. For the variables swelling power, paste clarity, and syneresis, data were adjusted to a regression model using the Sisvar software (Ferreira, 2011).

**Results and Discussion**

The starch extraction yield of *X. riedelianum* was 7.89%, similar to that of 6.61 to 11.81% found for arrowroot (*Maranta arundinacea* L.), a rhizomatous species also known as an unconventional vegetable (Souza et al., 2019). However, these extraction yields are much lower than that of 33.5%, expressed on a dry basis and considered highly efficient, obtained in the Brazilian industry of cassava (*Manihot esculenta* Crantz) starch (Branco et al., 2020). Therefore, there is a need to evaluate other methods to improve *X. riedelianum* starch extraction efficiency.

The average starch humidity level was 10.57±0.15 g 100 g⁻¹, close to that of 8.01 g 100 g⁻¹ found by Ávila et al. (2012) when also evaluating starch isolated from *X. riedelianum* rhizomes. However, these values are below the limits established by the Brazilian legislation for the good conservation of commercial starches: maximum of 21.0 g 100 g⁻¹ for potato (*Solanum tuberosum* L.) and maximum of 18.0 g 100 g⁻¹ for cassava (*Manihot esculenta* Crantz) (Anvisa, 2005).

Birefringent structures called Maltese crosses were identified under light microscopy (Figure 1 A). These structures are characteristic of unfused starch granules and are visible under polarized light due to the high degree of the supramolecular organization of the granules (Xiao et al., 2020). This means that the starch from *X. riedelianum* can be heated to 45°C without changing the configuration of the amyllose and amylopectin molecules, which could alter the functionality of the starch. This result is also indicative of the wide range of the gelatinization temperature of *X. riedelianum* starch. Therefore, for processes where gelatinization is not desirable, the drying temperature at 45°C is safe and, for processes where gelatinization is desired, higher temperatures are required (Alcázar-Alay & Meireles, 2015; Sjöö & Nilsson, 2018).

The particle size distribution of *X. riedelianum* starch is polymodal, with three peaks (Figure 2 A): the smallest at 1.20 μm, followed by the main one at 9.25 μm, and the third at 92.05 μm. The average volumetric diameter of the particles was expressed.
as the De Brouckere mean diameter [D (4.3)]. For \textit{X. riedelianum} starch, the granule diameter found was of 27.20 μm, close to that of arrowroot, with most granules with 20–35 μm in diameter (Souza et al., 2019). Granule size is important in numerous applications, influencing the reaction surface and settling speed during industrial processes. Small granules of 2 μm, for example, can be used as fat substitutes because their size is similar to that of lipid mycelia (Daiuto & Cereda, 2003). Larger granules, as in the case of those of \textit{X. riedelianum} starch, can be used in the production of biodegradable plastic films, as well as for bakery products (Alcázar-Alay & Meireles, 2015; Sjöö & Nilsson, 2018). Mango (\textit{Mangifera indica} L.) stick starch, because of its similar characteristics and granule size, can be used to prepare soups, broths, and porridges (Sjöö & Nilsson, 2018).

The microscopic analysis showed that the granules presented a smooth surface without superficial porosity and a spherical or semi-spherical shape with an irregular base (Figure 1 B), similar to that observed for cassava starch granules by Fernandes et al. (2019). Several compound starch granules were also found, which can be attributed to the presence of other residual compounds, such as proteins and lipids, as reported by Ashogbon & Akitayo (2012).

The morphology of the starch granules can interfere with the properties of the paste, the solubility of the starch, and the susceptibility of the granules to enzymatic action. The smooth surface of the granule observed in the present study hinders the access of enzymes and delays or hinders enzymatic action, increasing resistance to degradation (Vanier et al., 2019).

Starch is considered resistant when it is partially digested or undigested by the human digestive system, being widely used in the food industry as a dietary fiber, which is desirable for many food products and can be added to formulations. However, the evaluated starch is not the most suitable for the production of simple sugars due to its resistance to degradation by amylolytic enzymes.

Regarding diffraction, the X-ray diffractogram showed a lower intensity peak near 5.6° in \textit{X. riedelianum} starch, followed by more intense peaks when the 2Θ angle was near 17, 22, and 26° (Figure 2 B). Therefore, this starch exhibits a B-type diffraction pattern. Most root and tuber starches, such as those from the African arrowroot and different yam species, follow a similar pattern (Hoover, 2001).

The crystallinity index, determined from the total areas and peaks of the X-ray diffractograms, is an important parameter that influences the physical, mechanical, and technological properties of starch. The crystallinity found for \textit{X. riedelianum} starch was of 41.20%, considered high, but within the range suggested by Miranda et al. (2019), between 15 and 45% for native starches. Higher values like the one

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Morphology of \textit{Xanthosoma riedelianum} starch under optical microscopy using 40 x zoom (A) and micrography of the granules of the starch through scanning electron microscopy (B).}
\end{figure}
obtained in the present study imply greater granular stability, which, consequently, reduces the swelling capacity of the granule, making it more resistant to the gelatinization process (Singh et al., 2004). These results confirm that *X. riedelianum* starch is not easily gelatinized. For the food industry, this means higher temperatures and a greater energy expenditure for this process, indicating that chemical gelatinization may be more viable (Alcázar-Alay & Meireles, 2015; Sjöö & Nilsson, 2018).

For gelatinization, the initial temperature was 67.63±0.44°C, with a peak at 74.24±0.50°C and conclusion at 80.90±0.24°C; the temperature range was 12.61±0.68°C and enthalpy change was -4.94±0.078 J g⁻¹. If the maximum values of the standard deviations and variation are considered, the gelatinization of starch can occur at temperatures from 67.37 to 94.43°C; therefore, the obtained average value of 80.90°C represents a very high temperature.

Three thermal events of mass loss were observed for TGA/differential thermal analysis (Table 1). The first one corresponds to a mass loss of 9.66% and was attributed to the evaporation of volatiles, mostly of the water absorbed by the starchy material, which occurred almost at 92.72°C. The second event is associated with the thermal degradation phase of the major constituents of the starch and was composed by two peaks, beginning at 257.55°C and topping out at the degradation maximum of 326.38°C; the first peak was minor and is related to the loss of amylose, the most linear structure of starch, and the second refers to the degradation of the amylpectin structure, with a total mass loss of 75.74%. Finally, the third event is characterized by a small mass loss of 11.46% between 434.56 and 485.95°C due to the degradation of the carbonaceous material more resistant to temperature. This means industrial processes with temperatures above 92.72 and 257.55°C can destroy the structure of amylose and amylpectin, respectively (Alcázar-Alay & Meireles, 2015). When food is baked, for example, the temperature exceeds 250°C, although it may be slightly lower inside the food. In processes that include pressure increase, the temperature may be slightly lower, causing no damage to amylpectin, but degrading amylose (Sjöö & Nilsson, 2018). This can be positive for products in which amylose rearrangement is not desired, i.e., in which starch retrogradation is not the aim.

After the thermogravimetric analysis, it is possible to determine the level of inorganic substances in the sample.
studied samples (Sambale et al., 2019). In the present study, the burning residue represented 3.14% of the total starch mass; this small quantity shows that the starch was well extracted. According to Sjöö & Nilsson (2018), starch also contains a certain amount of minerals.

When analyzing paste properties, in the first 3.2 min (paste time), the starch granules did not change and viscosity was low, of 21±1.41 cp (Figure 2 C). Granules started to swell up at 77.9°C, called paste temperature, indicating that the starch has a high resistance to expansion and rupture, according to Singh et al. (2004). The observed temperature was similar to the one of 82.15°C reported by Ávila et al. (2012) for starch from *X. riedelianum* rhizomes. Therefore, the starch from this species needs a relatively high temperature to be gelatinized, requiring a greater energy expenditure and longer cooking time. For corn (*Zea mays* L.), the gelatinization temperature is, on average, 72°C (Sjöö & Nilsson, 2018).

After the beginning of paste formation, the granules started to swell, increasing viscosity to 3,263±7.07 cp, considered the peak in the heating cycle, reached in 4.13 min, at 88°C. During the heating period, the viscosity profile indicated some resistance of the bond forces inside the granules, which, when broken, under continuous agitation, caused a decrease of 1,016.5±6.3 cp in viscosity, that is, a break. Viscosity breakdown indicates the degree of granule disintegration after swelling and is calculated by the difference between peak and minimum viscosity. The lower the breakdown value, the greater the shear strength and stability of starch in processes involving heating (Andrabi et al., 2014).

The retrogradation tendency of *X. riedelianum* starch was of 970±5 cp, lower than that of 1,380 cp observed for cassava and higher than that of 804 cp for potato and of 828 cp for *Arracacia xanthorrhiza* Bancroft (Shirai et al., 2007). Alcázar-Alay & Meireles (2015) concluded that the higher the retrogradation tendency, the greater the diameter of the granules due to a higher fragility. In the industries that use starch and also in food industries, the most commonly marketed starches are those from corn, cassava, and potato due to their easy extraction and yield (Souza et al., 2019). In relation to tendency to retrograde, the studied *X. riedelianum* starch has an intermediate behavior to that of the corn and potato starches. When retrogradation is desired, the retrograded corn starch is more easily used in food products, such as chilled desserts; however, for products for which a lower consistency is desired, potato starch is preferred (Sjöö & Nilsson, 2018). Therefore, it is assumed that *X. riedelianum* starch can be well used in processes in which retrogradation does not occur so easily, specifically in products for which the lowest viscosity is desired during and after cooling.

The final viscosity of *X. riedelianum* starch was 3,116.5±19.09 cp, being higher than the values of 3,012, 2,448, and 2,520 found for potato, *A. xanthorrhiza*, and cassava, respectively, by Shirai et al. (2007). Final viscosity is an important parameter for the use of starch in food since it corresponds to the viscosity of a product at a given temperature; in this case, at 50°C. According to the type of product, a higher or lower viscosity can be desirable; instant soups, for example, must not present a very high final viscosity, which would cause an unpleasant sensation, whereas pie fillings require a higher viscosity to avoid its contents from overflowing during transportation (Garcia et al., 2014). Due to its high levels of viscosity, the starch from *X. riedelianum* would be better indicated for pie fillings, puddings, children’s food, and baking products. Daiuto & Cereda (2003) added that, for the calculation of the desired viscosity, it is necessary to consider the reduction that occurs in viscosity during industrial processing.

The swelling power of *X. riedelianum* starch does not change significantly between 25 and 70°C, but increases from 70 to 80°C, continuing up to 90°C (9.80±0.13 g); although the results suggest a plateau, there were no analyzes at higher temperatures (Figure 3 A). For food

| Phase | Mass loss (%) | Accumulative loss (%) | ΔT (°C) | Tp (°C) |
|-------|--------------|-----------------------|---------|---------|
| First | 9.66         | 9.66                  | 44.53–164.92 | 92.72   |
| Stability | -            | 9.66                  | 164.92–257.55 | -       |
| Second | 75.74        | 85.40                 | 257.55–434.56 | 326.38  |
| Third | 11.46        | 96.86                 | 434.56–485.95 | 439.35  |
| Residue | 3.14         | -                    | 485.95–600.00 | -       |

(1) ΔT, temperature variation; and Tp, peak temperature.
products, this means that *X. riedelianum* starch must be used in processes that require higher temperatures, above 70°C, in order for it to absorb more water and become more swollen. Starch swelling is one of the premises for gelatinization to occur (Alcázar-Alay & Meireles, 2015; Sjöö & Nilsson, 2018).

Singh et al. (2004) attributed the low swelling power of starch granules to the presence of several crystals formed by the association of long amylopectin chains, or, to a higher degree of crystallinity, which implicates in a higher stability and, consequently, decreases swelling capacity. In the present study, a high level of crystallinity of 41.20% was found for *X. riedelianum* starch using the X-ray diffraction analysis.

The average values and standard deviation of the solubility index are presented in a graph (Figure 3 B); however, the results did not fit to the proposed mathematical models. The behavior of *X. riedelianum* starch showed a constant growth of 1.18 to 1.85 g from room temperature to 70°C, followed by a drastic decrease to 0.66±0.22 g at 80°C, increasing again at 90°C. Although the graph is not obeying any mathematical equation, which could predict the behavior of the loss of solids by *X. riedelianum* starch yields as a function of temperature, a good observation and comparison between this graph and the one of swelling power shows that these two forces are complementary or even mirrored. When one value increases, the other decreases, which is the behavior expected for these two forces: the more a granule of common starch absorbs water, the more that granule loses solids (Sjöö & Nilsson, 2018).

The obtained results indicated once again that, for the food industry, the processes that involve cooking of *X. riedelianum* starch require higher temperatures, which translates in more time and a greater energy

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**Figure 3.** Regression of the swelling power of *Xanthosoma riedelianum* starch in function of temperature (A); solubility index of the starch as affected by temperature (B); percentage of the absorbance and transmittance of the starch in function of storage time (C); and effect of storage time over the syneresis of the starch (D).
expenditure when a greater swelling is desired for starch gelatinization, with a consequent increase in the loss of solids by the starch granules. However, when total gelatinization is not desired, *X. riedelianum* starch is a good option because, when added to food formulations, even under a heating process, it will not swell as much and will not reach the gelatinization stage if the processing temperature is below 70–80°C.

The percentage of absorbance and transmittance determines the clarity of the starch paste: the higher the transmittance, the greater the paste clarity (lower turbidity). On the first day, the transmittance of *X. riedelianum* starch presented the highest average of 4.88%, which decreased over time, reaching an average of 0.36% on the eighth day. The behavior of absorbance was opposite to that observed for transmittance (Figure 3 C). Hernández-Medina et al. (2008) reported 13.57 and 7.0% transmittance for arrowroot and taro (*Colocasia esculenta* (L.) Schott) starch, respectively, whereas Macena et al. (2020) found a value of 2.20% for avocado (*Persea americana* Mill.) seed starch.

Starches used as a thickener in pie fillings, in food toppings, or as edible films must be preferentially transparent, while starches used in the bakery, concentrated-drink, and processed-meat sectors may have the characteristics of the analyzed starch, as a clear paste is not necessary to formulate the products (Hernández-Medina et al., 2008; Egharevba, 2020). Therefore, the starch from *X. riedelianum* should be used in foods where translucency is not so important, as in fillings, flans, and puddings, or in the formulation of cakes, breads, and pies, and other bakery products.

The amount of water released by the *X. riedelianum* starch gels during storage increased up to 120 hours, when there was a decrease (Figure 3 D). For potato starch after 48 hours of storage, Colussi et al. (2018) verified water loss, or syneresis, of 46.10%, which is higher than that of 26.75±2.71% found for *X. riedelianum* starch during the same storage period. These different syneresis characteristics can be related to the differences in the quality of the recrystallized amylopectin crystals (Freschi et al., 2014). Water loss is associated with the reorganization of the molecules after cooling and storage, which can affect the functional properties of the starch in terms of viscosity and gel behavior (Charles et al., 2016), and is not considered beneficial to the food industry since it can affect the quality and lifespan of the food.

**Conclusions**

1. *Xanthosoma riedelianum* starch has a low extraction yield.

2. The morphology of *X. riedelianum* starch shows granules without surface porosity, resulting in a low susceptibility to enzymatic action.

3. *Xanthosoma riedelianum* starch requires high temperature and a longer cooking time to swell and achieve complete gelatinization.

4. *Xanthosoma riedelianum* starch is indicated for products that require high viscosity such as pie fillings, puddings, children’s food, and baking products, and its opaque and viscous paste can also be used to formulate broths.

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