Characterization of Calcium Carbonate Obtained from Oyster and Mussel Shells and Incorporation in Polypropylene

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There is a high content of calcium carbonate in mussel and oyster shells, which can be used in the formulation of medicine, in construction or as filler in polymer materials. This work has as its main objective to obtain calcium carbonate from mussel and oyster shells and used as filler in polypropylene compared their properties with polypropylene and commercial calcium carbonate composites. The shellfish was milling and heated at 500 °C for 2 hours. The powder obtained from shellfish were characterized by scanning electron microscopy (SEM), X-ray fluorescence, particle size distribution and abrasiveness and compared with commercial CaCO₃ and mixed with polypropylene. The thermal and mechanical properties of polypropylene with CaCO₃ obtained from oyster and mussel shells and with commercial CaCO₃ were analysed. The results showed that CaCO₃ can be obtained from oyster and mussel shell and is technically possible to replace the commercial CaCO₃ for that obtained from the shells of shellfish in polypropylene composites.

**Keywords:** oyster shell, mussel shell, calcium carbonate, polymer

1. Introduction

Aquaculture is an important source of animal protein for human consumption. In South America, Brazil is second in production of shellfish, following Chile¹. The state of Santa Catarina is responsible for more than 90% of Brazilian production of cultivated oysters and mussels. Bivalve shellfish production consists in the cultivation of the Mitilidae family represented by the perna perna mussel and the Ostreidae family, represented by the pacific oyster Crassostrea gigas².

After harvest, the product for the market (the shellfish meat) can go through a beneficiation process to improve its appearance and add value. In these units, the mussels are cooked by steaming or immersion and the shell is removed. Of the entire amount of shellfish produced, 75 to 90% consists of shells. These shells are composed of 95% calcium carbonate, and the remainder is organic matter and other compounds¹. The mussel and oyster shells are discarded into the environment. The improper disposal of solid waste from shellfish cultivation decreases water oxygen and microalgae that are responsible for the nutrition of mussels and oysters, thus hindering the growth of these shellfish¹.

Concern about the destination of oyster and mussel waste is not only observed in Brazil. In Korea, for example about 300,000 t of oyster shells are generated annually. The Korean government, concerned with public health, financed a project to increase recycling of this waste, because if this waste has been left untreated for a long time, it can be a source of nasty smell as a consequence of the decay of flesh remnants attached to the oyster or the microbial decomposition of salts into gases such as NH₃, H₂S and amines³. Many studies have investigated the use of oyster-shell waste as construction materials⁴⁵. Another possibility for reuse of these shells is to use them as supplementary feeding⁶⁷. Lee and co-workers² demonstrated that crushed oyster shells could be used as an alternative liming material to restore soil chemical and microbial properties in upland soil and to increase crop productivity.

Calcium carbonate is the most widely used filler in terms of the number of applications in polymer. The cheapest grades are low in price and are used primarily to reduce costs. By contrast, the finest grades are an order of magnitude higher in price and are used to modify various properties, both during processing and in the final compounds⁸. Some studies have investigated the use of oyster-shell waste as filler in polymer⁹⁻¹⁰. Chong et al.¹¹ prepared mixtures of recycled polyethylene (PE) and oyster-shell powder, that was coated with ionic surfactant, to test their fire-retardant properties. The mixtures present better mechanical properties and fire-retardant behavior compared with neat recycled PE. Funabashi et al.¹² evaluated method of biomass carbon ration of polymer filled with calcium carbonate using poly(butylene succinate) (PBS) with oyster shell powder and poly (lactic acid) (PLA) with no-biobased inorganic calcium carbonate. They observed that the estimation method is effective for polymer composites with CaCO₃.

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The aim of this work is to obtain calcium carbonate from seafood (mussels and oyster) shells and demonstrate that this material can be reused as filler in polymer, based on thermal, chemical and physical properties of calcium carbonate and mechanical and thermal properties of polypropylene/calcium carbonate composite.

2. Materials and Methods

The polypropylene (PP) and commercial calcium carbonate used in this work were supplied by Braskem and Crenor Carbonatos, respectively, under the tradename PP H604 and Crenor MAX EB respectively. To obtain calcium carbonate, mussel and oyster shells supplied by a restaurant in Joinville, Brazil were used.

The mussel and oyster shells were heated in an oven at 200 °C for 1 hour to make the shells more brittle and submitted to milling in a high-speed planetary mill with a porcelain jar and alumina balls for 15 minutes with water. The powders are heated again to 500 °C and maintained for 2 hours and to undo the clusters a new milling was performed without water for 1 minute. The powders were characterized by particle size distribution analyses and chemical composition using a laser diffraction analyzer (CILAS 1064) and X-ray fluorescence (SHIMADZU EDX-700).

Abrasiveness was analyzed using 100 g of each sample dissolved in distilled water and transferred into the test cup. This cup was connected to a PVC roller which carries 174,000 rounds on the solution in contact with a grille weighed in advance. After the test, the grille was weighed again and the abrasiveness was determined by calculating the mass difference before and after the test, relating to the area subject to abrasion. This technique was performed in an Einlehner AT 1000 device.

Polypropylene composite with commercial CaCO₃ and mussel and oyster shells were prepared using a single screw extruder (Ciola B50) set to a length/diameter equal to 25/1, screw rotation rate of 150 rpm, with a flat temperature profile along the barrel (180 °C). The composite composition studied was 10 wt. % of CaCO₃, and 90 wt. % of neat PP. The composites were injected into a Sandretto Micro 65 injection-molding machine to obtain specimens for tensile and impact tests. The injection molding was carried out with a temperature in the nozzle of 200-225 °C.

The thermal degradation of the calcium carbonate and composites was analyzed in a TA Instruments TG Q200 thermogravimetric analyser in a nitrogen atmosphere. Non-isothermal analysis was performed in a temperature range of 25-800 °C, at 10 °C/min and nitrogen flow was maintained at 50 cm³/min.

Charpy impact tests, following ISO 179, were performed on an EMIC Impact tester. Tensile tests, according to ISO 527 standards, were carried out in an EMIC universal testing machine model. Crosshead displacement rate of 10 mm/min was used. A longitudinal strain gauge was used to obtain the Young modulus.

The composites were analyzed by differential scanning calorimetry (DSC) to determine the melting temperature, percent of crystallinity and oxidation induction time (OIT). The DSC tests were performed on equipment from TA Instruments DSC Q20. The OIT analysis was conducting according to ASTM D 3895-98. For determination of melting temperature and percent of crystallinity, the samples were heated at 10 °C/min under nitrogen atmosphere. The percent of crystallinity (Xc) was determined from the enthalpy of crystallization of PP, Equation 1, using a value of ΔHf° = 165 J.g⁻¹ for isotactic PP 100% crystalline and the enthalpy values were corrected for PP weight present in the composite.

\[ X_c = \frac{\Delta H_{amoastra}}{\Delta H_f^0} \times 100 \]  

(1)

Changes in the morphology in the fracture surface during the impact test of the PP/CaCO₃ and CaCO₃ morphology were observed in scanning electron microscopy (SEM) Zeiss DSM. The samples were first coated with a thin gold layer.

3. Results and Discussion

Table 1 presents the chemical composition of commercial CaCO₃ and mussel and oyster shells. Mussel and oyster shells present a slightly lower, although not significant, amount of calcium oxide (CaO) than commercial calcium carbonate. There are differences in chemical composition because oysters and mussels are water filterer⁸. Kurunczi and co-workers⁹, for example, detected mercury (Hg) and lead (Pb) in mussel shells, because the water where these mussels grew was contaminated. Chou et al.⁹ also detected Pb in oyster shells in New Orleans. Mussel shells also present a higher quantity of Fe₂O₃ than oyster shells, this oxide can influence polymer oxidation when these shells are used as filler in polymer matrix.

Figure 1 presents commercial CaCO₃, mussel and oyster shell particle size distribution. It is possible to observe a larger particle size distribution to oyster and mussel shells (Figure 1a,b respectively) than commercial CaCO₃. This behavior can be associated with a higher hardness of the shells, because there is a higher concentration of silica in the shells that increases shell hardness²⁰ and the milling conditions used was kept constant. Commercial calcium carbonate grinding consists in several stages with different grinding systems, until it reaches the average particle size most appropriate to use¹³.

The MEV photomicrographs for commercial CaCO₃, mussel and oyster shell are shown in Figure 2. It is observed that both the oyster and mussel shells (Figure 2a,b, respectively).
respectively) contain higher concentration of fine particles and there is also some presence of larger particles which lead to more heterogeneous distribution compared to the commercial CaCO$_3$ (Figure 2c). These results are also being observed in particle size distribution curves of Figure 1. The shape of the oyster and mussel shells particles is more irregular than commercial CaCO$_3$.

The abrasiveness test showed that the commercial CaCO$_3$, mussel and oyster shells were 32, 28, 32 g.cm$^{-2}$ respectively, showing no significant variation in this property under the conditions tested.

Thermogravimetric analysis (TGA) and first derivative TG curve [(d(TG)/dT) denoted by DTG] of PP/CaCO$_3$ composites are shown in Figure 3a,b and the data are simplified in Table 2. Although the oyster and mussel shells contain other oxides in chemical composition, these did not influence significantly the behavior of thermal stability compared to the PP with commercial CaCO$_3$. Composites of PP with oyster or mussel presented higher temperature of onset decomposition (dO) and maximum of decomposition (doff).

Melting temperature ($T_m$), percent of crystallinity ($X_c$) and oxidation induction time (OIT) of PP/CaCO$_3$ composites are shown in Table 3. It can be noted that CaCO$_3$ obtained from mussel and oyster shells do not show any significant difference in melting temperature and percent of crystallinity compared to the composite with commercial CaCO$_3$.

Oxidation induction time is lower for the composites containing CaCO$_3$ obtained from oyster and mussel shells compared to the composite with commercial CaCO$_3$. This behavior can be explained by differences in chemical composition of the fillers. The mussels showed 0.715% Fe$_2$O$_3$, oyster shells less than 0.2% and in commercial CaCO$_3$, Fe$_2$O$_3$ was not detected. According Paoli$^{21}$ transition metal oxide (including iron) can catalyze polyolefin

Table 2. Degradation temperature of PP with oyster shell powder, mussel shell powder and commercial CaCO$_3$ from TG and DTG curves.

<table>
<thead>
<tr>
<th>Composites</th>
<th>dO (°C)</th>
<th>doff (°C)</th>
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<tbody>
<tr>
<td>PP/CaCO$_3$</td>
<td>256</td>
<td>401</td>
</tr>
<tr>
<td>PP/mussel shell</td>
<td>306</td>
<td>416</td>
</tr>
<tr>
<td>PP/oyster shell</td>
<td>292</td>
<td>421</td>
</tr>
</tbody>
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<table>
<thead>
<tr>
<th>Composites</th>
<th>$T_m$ (°C)</th>
<th>$X_c$ (%)</th>
<th>OIT (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PP/CaCO$_3$</td>
<td>159</td>
<td>45</td>
<td>84</td>
</tr>
<tr>
<td>PP/mussel</td>
<td>159</td>
<td>50</td>
<td>14</td>
</tr>
<tr>
<td>PP/oyster</td>
<td>160</td>
<td>46</td>
<td>45</td>
</tr>
</tbody>
</table>

Table 4. Mechanical properties of PP with oyster shell powder, mussel shell powder and commercial CaCO$_3$.

<table>
<thead>
<tr>
<th></th>
<th>PP/CaCO$_3$</th>
<th>PP/oyster</th>
<th>PP/mussel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Young's Modulus</td>
<td>918 (±58.1)</td>
<td>904 (±40.4)</td>
<td>902 (±49.6)</td>
</tr>
<tr>
<td>Yield strength</td>
<td>33 (±1.4)</td>
<td>35 (±0.6)</td>
<td>33 (±1.0)</td>
</tr>
<tr>
<td>Elongation of break</td>
<td>61 (±21.3)</td>
<td>38 (±18.2)</td>
<td>26 (±6.5)</td>
</tr>
<tr>
<td>Impact strenght</td>
<td>3.2 (±0.07)</td>
<td>3.4 (±0.42)</td>
<td>3.4 (±0.42)</td>
</tr>
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oxidation. Allen et al.\textsuperscript{22} conducted a test on samples of infrared PP containing silica which detected the presence of carbonyl, evidence of oxidation; the authors argue that the carbonyl formation is more associated with the presence of aluminum, titanium and iron than the presence of silica. Table 4 summarizes the results for mechanical property tests obtained for all samples studied. It is possible to observe that PP composites with commercial CaCO\textsubscript{3} and those obtained from shellfish do not show any significant difference on Young’s modulus, yield strength and impact

![Figure 2. SEM micrographs of CaCO\textsubscript{3} obtained from a) oyster and b) mussel shells; and c) commercial CaCO\textsubscript{3}.](image)

![Figure 3. a) TG curves and b) DTG curves of PP with oyster shell powder, mussel shell powder and commercial CaCO\textsubscript{3}.](image)

![Figure 4. SEM micrographs of impact test of PP with a) commercial CaCO3; b) mussel shells; and c) oyster shells.](image)
strength. The composite with commercial CaCO$_3$ showed higher elongation at break than those composites with fillers obtained from shellfish, which can be explained as being due to the particle size distribution.

Sample fractures resulting from impact experiments were analyzed by SEM, as shown in Figure 4. In Figure 4b,c that display the micrographs of the composites with mussel and oyster shells, we can see the large particles and many fine particles of CaCO$_3$ in the PP matrix, while the composite with the commercial carbonate presented particles smaller and more uniform. These differences can justify the smaller deformation at break of PP with shellfish composites. No adhesion between the domains in the matrix can be observed, probably due to the weak interfacial bonding in all composites.

4. Conclusion

The results of this study showed that CaCO$_3$ can be obtained from oyster and mussel shells, since both have a similar amount of CaO to commercial CaCO$_3$. The differences in particles size and distribution of particle size observed are due to the milling conditions used and firing conditions of calcinations that it was kept constant. Analyzing the tensile and impact test results, we can conclude that the composites with commercial CaCO$_3$ and mussel or oyster shells were similar, despite the great difference in particle size and distribution of particle size. For an amount of 10% load in the PP, there was no significant change in the mechanical properties, which makes this project technically feasible.

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References