Austenitic Stainless Steels
Mass Balance Mathematical Model
Carbon Dioxide Dynamic Adsorption
Artificial Oil Field Formation

Discovering Thoughts, Inventing Future
### Editorial Board

**Global Journal of Research in Engineering**

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Evaluation of the Corrosion Behavior on Austenitic Stainless Steels in Artificial Oil Field Formation Water using Potentiodynamic and Potentiostatic Electrochemical Techniques

By Jorge Luiz Cardoso, Luís Flávio Gaspar Herculano, Pedro de Lima Neto & Marcelo José Gomes da Silva

Federal University of Ceará

Abstract- Objective: The main objective of this research was to evaluate the corrosion resistance of some austenitic and super austenitic stainless steel when immersed in an aqueous solution of artificial oil field formation water saturated with CO₂ and also without CO₂. With these two simulated situations, the pH effect of the solution on the corrosion resistance of the alloys was studied.

Methods: Samples of austenitic and super austenitic stainless steels previously characterized by x-ray diffraction using synchrotron light were used in this research. Two electrochemical techniques, one potentiodynamic and the other potentiostatic were used to investigate the effect of the solution on the corrosion behavior of the samples.

Keywords: austenitic stainless steels, CO₂ corrosion resistance, aqueous solution, pitting corrosion, potential step, passive layer.

GJRE-C Classification: NLM: WC 800

Strictly as per the compliance and regulations of:
Evaluation of the Corrosion Behavior on Austenitic Stainless Steels in Artificial Oil Field Formation Water using Potentiodynamic and Potentiostatic Electrochemical Techniques

Jorge Luiz Cardoso ♦, Luís Flávio Gaspar Herculano ♦, Pedro de Lima Neto ♦ & Marcelo José Gomes da Silva ♦

Abstract: Objective: The main objective of this research was to evaluate the corrosion resistance of some austenitic and super austenitic stainless steel when immersed in an aqueous solution of artificial oil field formation water saturated with CO₂ and also without CO₂. With these two simulated situations, the pH effect of the solution on the corrosion resistance of the alloys was studied.

Methods: Samples of austenitic and super austenitic stainless steels previously characterized by x-ray diffraction using synchrotron light were used in this research. Two electrochemical techniques, one potentiodynamic and the other potentiostatic were used to investigate the effect of the solution on the corrosion behavior of the samples. The linear polarization test was used to evaluate the CO₂ corrosion resistance of the alloys in artificial oil field formation water. The potentiostatic technique (potential step) was used to assess the influence of the solution without CO₂ on the corrosion resistance of the alloys, thus, varying the pH of the electrolyte. Scanning Electron Microscopy (SEM) was used to observe the type of corrosion on the surface of the samples.

Results: The results indicated that the type of corrosion found on the surface of the alloys was pitting corrosion. The pH effect of the solution (artificial oil field formation water) influenced the pits' shape. The conventional austenitic steels showed to have low corrosion resistance in chloride-containing environment. The super austenitic stainless steels presented a high corrosion resistance in the solution with and without CO₂. No pits or micro pits were observed on their surfaces by SEM.

Conclusion: From the results obtained, it was evident that the conventional austenitic stainless steels are not a good choice in severe environments like those found in the pre-salt region. The super austenitic stainless steels showed to be a good option in CO₂-containing environments, mainly in aqueous solution with a high content of chloride.

Keywords: austenitic stainless steels, CO₂ corrosion resistance, aqueous solution, pitting corrosion, potential step, passive layer.

I. Introduction

The discoveries made in the pre-salt region (a geological formation of continental shelves) in Brazil are among the world’s most important in the past decade. In this region, there is a considerable amount of good quality oil and this reality puts Brazil in a strategic position for the global demand for energy [1]. The discovery of this region brings several technological challenges for the oil and gas exploration. The corrosion process in this region occurs under specific conditions. Some of them are high temperatures between 80°C and 150°C, the presence of gases such as carbon dioxide (CO₂) and hydrogen sulfide (H₂S), oil formation water, high pressure, leaving the operating environment very hostile[2]. The main characteristic of the pre-salt region is the high content of sodium chloride (NaCl) found there. This NaCl, CO₂ and H₂S dissolved in the oil field formation water can accelerate the corrosion of metallic materials used for the oil exploration in the pre-salt region. Another concern for this operation is the environmental impact that can occur if these materials fail. Cheaper materials such as carbon steels are a good choice but there is a problem related to them: The difficulty in adding corrosion inhibitors for carbon steel pipes in offshore oil extraction at great depths. This has led to the increased use of corrosion resistant alloys[3]. Of all types of corrosion, localized corrosion, especially pitting corrosion, is the most common in marine waters and difficult to control. Currently, the oil and gas industry is concerned about the environmental impact caused by oil leaks in the marine ecosystem. This type of accident can be prevented using materials more resistant to the environmental conditions found in the pre-salt region. There are two types of technological challenges for the exploration of oil and gas contained in the pre-salt region: the other challenge consists of drilling the well as far as the reservoir, crossing water layers, sediment, and salt. Each layer with a different behavior at temperatures ranges from 50°C to 150°C under high pressures and corrosive gases, all these conditions acting together. The way back to the surface must also be considered.
All the oil and natural gas extracted from the well will be transported through the pipelines, and the material from which the pipes are made must resist all adverse conditions to avoid oil leaks. The second challenge is horizontal and consists of transporting the oil and gas from the production area to the coast, localized about 300 km away from the well location [1]. In summary, it is a set of problems that begins with the well’s depth, passing by the coating when drilling into soft sediments through the salt layer to reach a very high temperature and pressure environment saturated with corrosive gases already mentioned [4]. Corrosion resistant-alloys such as super austenitic stainless steels are a great choice when considering the severe conditions of pre-salt. Due to their high chromium, nickel, and molybdenum content, it is expected that this kind of material presents more corrosion resistance in the pre-salt conditions than the conventional stainless steels. Some authors have already studied the corrosion resistance of austenitic stainless steel and other materials regarding the effect of CO₂ [1,5–8]. For this paper, the effect of the oil field formation water with and without CO₂ was studied, taking into account two austenitic and two super austenitic stainless steels for later comparison. Two electrochemical techniques, one potentiodynamic and the other potentiostat were used to evaluate the corrosion resistance for these materials in artificial oil formation water.

a) Materials

For this research, the materials used were the AL-6XN PLUS™ super austenitic stainless steel, the 904L super austenitic stainless steel, and the 300 series austenitic stainless steels AISI 316L and 317L. The chemical composition of the materials studied presented in Table 1 were measured in an Optical Emission Spectrometer (PDA-7000 SHIMADZU). The Pitting Resistance Equivalent Number (PREₙ) was calculated using equation 1.

\[
\text{PRE}_n = \%\text{Cr} + 3.3\ \text{Mo} + 30\%\text{N} \quad (\text{Eq.1})
\]

Table 1: Chemical composition (wt%) of the studied alloys and the respective Pitting Resistance Equivalent Number (PREₙ)

<table>
<thead>
<tr>
<th>Alloys</th>
<th>C</th>
<th>N</th>
<th>Mn</th>
<th>Si</th>
<th>Cr</th>
<th>Ni</th>
<th>Mo</th>
<th>PREₙ</th>
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<tr>
<td>316L</td>
<td>0.030</td>
<td>0.05</td>
<td>1.65</td>
<td>0.41</td>
<td>17.2</td>
<td>10.7</td>
<td>2.2</td>
<td>26</td>
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<tr>
<td>317L</td>
<td>0.024</td>
<td>0.06</td>
<td>1.49</td>
<td>0.40</td>
<td>17.8</td>
<td>12.3</td>
<td>3.5</td>
<td>31</td>
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<tr>
<td>904L</td>
<td>0.027</td>
<td>0.10</td>
<td>0.74</td>
<td>0.66</td>
<td>19.5</td>
<td>24.3</td>
<td>4.5</td>
<td>37</td>
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<tr>
<td>AL-6XN PLUS™</td>
<td>0.021</td>
<td>0.24</td>
<td>0.35</td>
<td>0.32</td>
<td>21.8</td>
<td>25.8</td>
<td>7.6</td>
<td>54</td>
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b) Characterization of the Samples

The samples in the as-received condition were characterized using x-ray diffraction (XRD) by Synchrotron Light (energy 12 keV) to detect the phases. ICDD database (International Centre for Diffraction Data) was used to identify the peaks of the phases. A Gleeble was used to fix the samples. The measurements were carried out at the Brazilian Synchrotron Light Laboratory in the city of Campinas-SP in Brazil. For this characterization, the shape and dimensions of the samples are shown in Figure 1.

c) Electrochemical Tests used

To evaluate the corrosion behavior of the materials, two electrochemical techniques were used: a potentiodynamic technique (linear polarization test) and a potentiostatic technique (potential stepstest), one complementing the other. Firstly, the linear polarization technique was used to evaluate the effect of CO₂ on the corrosion behavior of the samples. The dimensions of the samples were 5.0 mm x 5.0 mm x 3.5 mm with an average exposure area of 39 mm². The samples were mounted in cold resin and ground using SiC paper up to 600 mesh, washed in distilled water, and then blow-dried. An adapted cell was used with two gas inlets, one for CO₂ and one for N₂. The cell also contained an input for a pH reader, in addition to the classic inputs for the three main electrodes (reference, working, and counter electrodes) in addition to a gas outlet (see Figure 2). The electrodes used were the samples (working electrode), as counter electrode a platinum electrode (93 mm²), and the reference electrode used was the silver chloride silver (Ag/AgCl/Cl-sat) saturated with KCl. The electrolyte used (artificial oil field formation water) was named by Petrobras of TQ 3219 which composition is shown in Table 2.

Table 2: Chemical composition of the artificial oil field formation water for 1 L of distilled water

<table>
<thead>
<tr>
<th>Reagents</th>
<th>CaSO₄</th>
<th>MgCl₂</th>
<th>NaHCO₃</th>
<th>NaCl</th>
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<tr>
<td>C (g/L)</td>
<td>0.516</td>
<td>4.566</td>
<td>0.425</td>
<td>29</td>
</tr>
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</table>

First, the solution was deaerated with N₂ gas to simulate the pre-salt environment (absence of free O₂) that could interfere with the results. The N₂ gas was bubbled into the solution until a pH of 8.2 ± 0.1. After this procedure, the solution was bubbled with CO₂ until saturation (pH 5.1 ± 0.1). In this procedure, the electrolyte became acidic. CO₂ gas in contact with an aqueous solution (oil field formation water) forms acids.
that react with the metallic elements of the alloy [9]. A potentiostat (AUTOLAB PGSTAT302N) connected to a microcomputer was used for both techniques. The software NOVA 1.9 was used to obtain data from the linear potential curves. Before the measurements, the samples were immersed for 30 min in the solution to determine the open circuit potential (OCP). The sweep of the polarization curves was -0.5 V to 1.2 V from the OCP with a sweep rate of 1 mV/s. After linear polarization tests, the samples were washed with water and sprayed with alcohol to clean the surface. Scanning Electron Microscopy (SEM) micrographs on the surfaces of the samples were obtained after corrosion tests for later comparison. The corrosion tests were reproduced in triplicate.

The potential step technique (potentiostatic technique) was also used to evaluate the corrosion behavior of the samples for the solution of artificial oil field formation water, this time with no CO₂ and no N₂. This test was intended to evaluate only the effect of the artificial oil field formation water on the surface of the samples. For this test, the samples were mounted in cold-curing epoxy resin, ground up to 600, rinsed with ethanol, and blow-dried before each measurement. The samples had the dimensions of 8.3 mm x 8.2 mm x 3.7 mm. To reduce crevice corrosion on the epoxy/steel, the specimens were coated with a lacquer leaving an exposed area of 1 cm². A three-electrode cell configuration was used. A saturated silver/silver chloride (Ag/AgCl) as reference electrode and a platinum electrode as a counter electrode were used. The electrolyte used was the same used in the linear polarization test (see Table 2). A potentiostat (AUTOLAB PGSTAT302N) connected to a microcomputer along with the software NOVA 1.9 was used. Before the measurements, the samples were immersed for 30 min in the solution to determine the open circuit potential (OCP), the same procedure used before. Subsequently, the potential was increased in steps of 50 mV every one hour until a breakthrough current density was attained. The pitting corrosion initiation potential was defined when the current density reached values above 0.1 mA/cm² [10]. After the tests, the samples were examined by SEM to confirm the presence of pits on their surfaces. The tests were carried out in triplicate at 25°C (room temperature).

II. Results and Discussion

a) Characterization of the Samples for the as-received Conditions

The X-ray diffractogram pattern for the 316L and AL-6XNPLUS™ steels can be seen in Figure 3. For both sheets of steels, the main phases detected were the matrix phase (austenite) and some ferrite peaks, indicating that both materials were not in the solution annealed condition. No other phases were detected for the analyzed angle range 2θ (25-79°). For this measurement, a synchrotron light radiation source (λ = 0.10332 nm) was used. This measurement was not possible for the 317L and 904L steels due to a manufacturing problem of the samples.

b) pH Study of the Solution

Firstly, the solution pH used in the corrosion tests (artificial oil field formation water) was studied. The solution was deaerated by bubbling N₂ to simulate the absence of free oxygen from the pre-salt layer. Figure 4 shows the results for the pH study of the solution (called TQ3219 by Petrobras). The stabilization of pH indicates that the electrolyte is deaerated and subsequently saturated with CO₂. All the chemical reaction that happens when bubbling CO₂ in the solution is described in our previous work [4].

c) CO₂ Corrosion Evaluation using a Potentiodynamic Technique

Before the linear polarization tests, the OCP of the samples was measured. The result is shown in Figure 05. The OCP of the super austenitic steels (AL-6XN PLUS™ and 904L) stabilize in 5 minutes. For the other austenitic steels (316L and 317L), the stabilization time is longer, especially for the 317L steel. After 30 minutes of immersion, all the OCP are stabilized. Figure 6 shows the linear polarization curves for the steels in the as-received condition. The linear polarization tests aim to verify the formation of passive film or not on the alloys surfaces[11]. For this test, all the samples were immersed in the solution used (artificial oil field formation water) saturated with CO₂. The super austenitic stainless steels AL 6XN PLUS™ and 904L showed a good CO₂ corrosion resistance. After reaching the corrosion potential (around -0.5 V), a passive film is formed and broken at -0.34 V until they reach a passivation peak around -0.20 V. After this potential, there is the formation of another passive layer that remains until the potential of +0.89 V where there is a slight breakdown of this layer and another passivation. After reaching a potential of +1.02V (pitting potential), there is an increase in current density, the transmissive region, since the potential is too high (above +1.0 V). It is possible to observe that the electrochemical behavior for the super austenitic steels studied in this work is very similar. Their passive regions are quite stable. The increase of current density after +1.0 V can be associated with oxygen evolution, reported in the literature[12]. The 317L steel also showed a good CO₂ corrosion resistance. The formation of its passive layer is not so stable as the passive layers of the super austenitic steels; even so, the current density in the passive region remains low, in the order of 10⁻⁶ A/cm². Its pitting potential is around +0.61 V. The 316L steel did not present any passivation since the anodic current increased with time. This steel presented the highest anodic current rate (in the order of 10⁻⁵ A/cm²) if
compared with the other steels. Its corrosion potential is similar to the corrosion potential of 317L steel (+0.40 V). 316L steel presented the lowest pitting potential (+0.52 V), indicating that its CO₂ corrosion resistance is not very effective. The reduction in anodic current density is associated with the passive film as a protective barrier against corrosion. The super austenitic steels AL-6XN PLUS™ and 904L showed a reduction in their anodic current, while the austenitic steels 316L and 317L showed an increase in anodic current with time. This result shows that the passive film of super austenitic steels is more stable. This effect can be attributed to the high levels of alloying elements such as Cr, Mo, and Ni. According to Sedriks, on a polarization curve, the greater the difference between the pitting potential and the corrosion potential (ΔE = Epit - Ecorr), the more resistant to corrosion the material is [13]. Table 2 shows the corrosion potential values, pitting potential, and the difference between them for the studied steels. The ΔE interval is higher for the super austenitic steels, which confirms their high performance about CO₂ corrosion. The 316L steel had the lowest value for ΔE, indicating that it is not a suitable material for applications that require good CO₂ corrosion resistance. More detailed work on CO₂ corrosion using austenitic stainless steels by the authors of this research can be found in [4].

Table 2: Potentials in V (Ag/AgCl) taken from the linear polarization curves for the studied steels

<table>
<thead>
<tr>
<th>Alloy</th>
<th>E(corr)</th>
<th>E(pit)</th>
<th>ΔE</th>
</tr>
</thead>
<tbody>
<tr>
<td>316L</td>
<td>-0.41</td>
<td>0.30</td>
<td>0.71</td>
</tr>
<tr>
<td>317L</td>
<td>-0.41</td>
<td>0.61</td>
<td>1.02</td>
</tr>
<tr>
<td>904L</td>
<td>-0.49</td>
<td>1.02</td>
<td>1.51</td>
</tr>
<tr>
<td>AL-6XN PLUS™</td>
<td>-0.53</td>
<td>0.99</td>
<td>1.52</td>
</tr>
</tbody>
</table>

After the linear polarization tests, SEM of the surfaces of the steels was carried out. The only material that presented pits on its surface was the 316L steel, as shown in Figure 7. Several factors may have influenced this form of corrosion for the 316L steel. Among them, one can mention: inefficiency of the passive film, pH of the solution, chloride content in the solution, effect of CO₂. The action of the chloride ion in an acid medium caused by the reaction of CO₂ gas in an aqueous medium can accelerate the localized corrosion process.

d) The Corrosion Resistance of the Steels in Artificial Oil Field Formation Water by Potential Step Technique

The samples in the as-received condition were submitted to another electrochemical technique called potential step. This time, no CO₂ or N₂ was used in the solution (artificial oil field formation water). The investigation using this technique is in agreement with the linear polarization experiments where the 904L and AL-6XN PLUS™ austenitic stainless steels had excellent pitting corrosion resistance when compared with the other austenitic steels (316L and 317L). Figure 8 shows the results of potential step for the 316L, 317L, 904L, and AL-6XN PLUS™ alloys, respectively. The graphs are of type double Y-axis, where the potential (V vs. Ag/AgCl, sat KCl) and the current density (mA/cm²) are plotted on the Y-axis, and the time (s) is plotted on the X-axis. Every potential step was maintained for one hour. If nothing happened on the passive film, then a new step was reached by an increment of +50 mV. The pitting potential (Eₚ) of each alloy was achieved when the current density reached values above 0.1 mA/cm², as shown on the graphs. So there was an abrupt increase in the current density indicating the breakdown of the passive film. The time to achieve the pitting potential depends on the film resistance of each alloy. The more resistant the passive film, the more time is needed to reach the pitting potential. The pitting potential for the 316L steel presented the lowest value (+0.52 V), while the pitting potential for the 904L and AL-6XN PLUS™ steels presented the highest value (+1.06 V and +1.09 V, respectively). The pitting potential for the 317L steel showed an intermediate value (+0.81 V). Table 3 shows the pitting potential and the time to achieve it for each alloy. It was necessary more than one day for the sample of the AL-6XN PLUS™ steel to reach its pitting potential. This result shows how resistant this material is to the conditions used. On the other hand, the 316L steel presented the lowest time to reach its pitting potential. Even without the presence of CO₂, this steel showed susceptibility to pitting corrosion in chloride-containing environments. The 317L steel showed to be more resistant than the 316L steel in chloride-containing environments but less resistant than the other two super austenitic steels. If compared with table 2, it can be seen that the pH of the solution shifted the pitting potential of the steels. In the presence of CO₂, the solution is more aggressive, decreasing the pitting potential of the steels.

Table 3: Measured pitting potential of the studied alloys using the Potential step technique

<table>
<thead>
<tr>
<th>Potential step</th>
<th>Alloy</th>
<th>E(pit) (V Ag/AgCl)</th>
<th>time (h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>316L</td>
<td>+0.52</td>
<td>13.5</td>
<td></td>
</tr>
<tr>
<td>317L</td>
<td>+0.81</td>
<td>16.2</td>
<td></td>
</tr>
<tr>
<td>904L</td>
<td>+1.06</td>
<td>23.1</td>
<td></td>
</tr>
<tr>
<td>AL-6XN PLUS™</td>
<td>+1.09</td>
<td>26.2</td>
<td></td>
</tr>
</tbody>
</table>

The 316L and 317L steels suffered pitting corrosion. For the 316L steel (Figure 9a), the pits possess a circular shape with a center hole. The pit propagates from the center to the edge and tries to grow with time. This effect is attributed to the chloride in the solution. The chloride ion (Cl⁻) is very small and can penetrate easily in sites of the 316L surface where the film is broken. The pits on the 316L steel grow but only in the center, as shown in Figure 9a. With the absence
of CO₂ in the solution, the environment is not so aggressive to permit the pits’ growth. The 317L steel also suffered pitting corrosion, but its pits are so small compared with the ones of the 316L steel. The pits initiated, but they did not grow with time, as shown in Figure 9b. These pits are non-uniform. This result indicated that the 317L steel in some chloride-containing environments is also resistant, being also a good choice in some applications where the 316L cannot be used, for example, in the oil and gas industry in chloride-containing environments. The super austenitic stainless steels (904L and AL-6XN PLUS™) presented pits much smaller than the ones found on the surface of conventional austenitic steels, as seen in Figure 9 (c-d). They are micro-pits, and after initiating, they passivate again before starting to grow. The effect of CO₂ on the morphology of the pits for the 316L steel, the most affected steel in the experiments, is shown in Figure 10. In Figure 10a, one can see the pits for the potentiodynamic test using CO₂ in the solution. In Figure 10b, it can be seen a single pit formed for the potentiostatic test with no CO₂ in the solution. For the potentiostatic test (potential step), the pit did not grow as expected, leaving a hole in the center. All the micrographs of the alloys taken after the corrosion tests are in accordance by the graphs shown before. This experimental procedure has previously been used to qualify Ni-based alloys and hyper duplex stainless steel for raw seawater injection [14] and also used to study a 13% Cr supermartensitic stainless steel related to localized corrosion [15]. These results combined with the linear polarization tests in CO₂-saturated aqueous solution, show that these materials (the super austenitic stainless steels) are an excellent option for chloride-containing environments with and without CO₂ once they are cheaper than the Ni-based alloys. In some cases, the conventional 317L steel can also be a good option than the conventional 316L steel.

III. Conclusions

It can be concluded that the oil field formation water plays an important role as an aggressive substance in the pre-salt region. In chloride-containing environments, the 316L steel is not so resistant, and it is not recommended for the content of NaCl in the pre-salt region. The type of corrosion found was identified as pitting corrosion. For the 316L steel, pits were formed for both techniques used, but the pitting corrosion was more aggressive for the potentiodynamic technique due to CO₂ in the solution and the absence of free oxygen. The 317L steel presented good pitting corrosion resistance when compared to the 316L steel. The two super austenitic stainless steels studied in this research (904L and AL-6XNPLUS™) presented good pitting corrosion resistance. Both can be the solution for applications in chloride-containing environments as those found in the pre-salt region.

Acknowledgments

The authors would like to thank to the Coordination for the Improvement of Higher Education Personnel (CAPES) and Cearense Foundation to Support Scientific and Technological Development (FUNCAP) for the financial support. A special thank is given to Wilman Italiano, who gave training in the electrochemical tests of potential step and also gave important contributions to this work.

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Adsorption Kinetics and Mass Balance Mathematical Model of Monoethanolamine Surface-Modified Palm Shell Activated Carbon for Carbon Dioxide Dynamic Adsorption in Fixed Bed Column

By Saad Hashim Khalil

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Keywords: adsorption kinetics, activated carbon, impregnation, MEA, fixed bed column.

GJRE-C Classification: LCC: TP156.A3

Strictly as per the compliance and regulations of:
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I. Introduction

The removal of carbon dioxide (CO₂) is significant for oil and gas industry due to its harmful acidic effects on oil and gas pipelines with no added energy value to natural gas [1]. Because of the increasing indications of CO₂ implication in global warming [2], capturing CO₂ from its emitting sources is becoming a vital topic. Adsorption is offering an effective alternative for CO₂ capturing comparing to other capture technologies [3]. There are many types of adsorbents; conventional, like, activated carbons, silica gel, ion-exchange resins, zeolites, and mesoporous silicates, activated alumina, metal oxides, and new like, carbon fibers and metal-organic frameworks. [4]. Adsorbent most important feature is adsorbing capacity [5], besides, good adsorbent should be selective and chemically and mechanically durable [6]. AC is cost-effective and adaptable microporous adsorbent [7] and is considered a superb adsorbent due to its high specific surface area, appropriate pore size distribution, diversity of surface chemistry [8]. It’s mostly micropore structure were used extensively in liquids and gases systems. The micropores and mesopores of the AC particles were utilized to accommodate the impregnating molecules, which can be attached chemically (grafting) or physically (impregnation) to the AC particles [9]. Impregnation of AC particles with chemicals improves their natural adsorption capability to adsorb gases [10]. Alkanolamines, such as, monoethanolamine (MEA), diethanolamine (DEA), and methyl diethanolamine (MDEA) are very important absorbents for acidic gases in the field of natural gas sweetening and for mitigation the adversity of these gases on environment [11] and they are extensively used in CO₂ absorption from different gas sources [12]. MEA, which is a primary amine, has been used intensively to capture CO₂ from gas streams and from many various sources due to its fast reaction kinetics with CO₂, low cost and thermal stability, as it is more favorable than other alkanolamines [13] [14]. Because of the effectiveness of liquid amine absorption process researchers were encouraged to utilize amines in their solid state for CO₂ capture [15]. Adsorption kinetics is essential tool used to evaluate the performance of an adsorbent and to understand the mechanism of adsorption [16] and many researches had included kinetics of batch CO₂ adsorption on different adsorbents in their works [17]. They found that the restriction step is the intraparticle diffusion (pore diffusion). On contrary to the findings of this paper where the restrictive step to CO₂ adsorption was the film diffusion. In this research, dynamic adsorption experiments were conducted to investigate the adsorption kinetics of MEA-impregnated AC particles packed in adsorption column to adsorb CO₂ from gas mixture.

II. Materials and Methodology

a) Materials

1. Certified analytical reagent monoethanolamine (MEA), C₂H₇NO, molecular weight 61.
2. Commercial palm shell AC was purchased from Bravo Green SDN BHD (Sarawak, Malaysia).
3. Gases
   a. Mixture of 15% CO₂ with 85% N₂.
   b. Pure N₂.

b) Methodology
   i. AC Particles Characterization
      Granulated palm shell AC particles were physically activated by steam. The mostly micropore particles have total Bet surface area of 838 cm²/g, while the micropore surface area of that total area is 675 cm²/g and micropore volume is 0.32 cm³/g.
   ii. AC Beds Perpetration
      A household coffee grinder crushed the AC particles. 710 and 500 µm sieves were employed to characterize the AC particles to the required particle size of 500 µm (particles passing 710 and stopping on 500 µm sieve).
   iii. Impregnation of AC Samples
      Impregnation was carried out by placing 5 g of granular AC in a beaker, 2 g of MEA added to the beaker with 10 g of deionized water as an environmentally friendly medium and to facilitate the impregnation process. The beaker contents were stirred at 500 rpm for 1 hour at room temperature. The final slurry then dried completely in Heraeus Instrument Vacuthermo oven at 70 °C under 0.1 bar vacuum pressure (absolute) for 6 hours. Samples of AC particles prepared for CO₂ adsorption separation experiments are, non-impregnated AC particles and MEA-impregnated AC particles.
   iv. Working Breakthrough Time
      Working breakthrough time was utilized as a method to evaluate the performance of AC beds.

\[
R_1R_2N^+HCOO^- + R_1R_2NH \leftrightarrow R_1R_2NCOO^- (\text{Carbamate}) + R_1R_2NH^+ \quad (1b)
\]

The overall reaction is as in equation 1:
\[
\text{CO}_2 + 2R_1R_2NH \leftrightarrow R_1R_2N^+HCOO^- + R_1R_2NH^+ (\text{Zwitterion}) \quad (1a)
\]

vi. Rate and Mechanism of Adsorption
   To investigate the rate of adsorption two equations were explored namely, pseudo first order (SFO) and pseudo second order (PSO) equations. The mechanism of adsorption and the adsorption controlling step was determined by Weber-Morris intraparticle diffusion model.

III. RESULTS AND DISCUSSIONS

a) Impacts of MEA Surface-Modification on AC Particles
   MEA molecules occupied the pores of the mostly micropore AC particles and blocked them, reducing significantly the micropore surface area from 675 to 36m²/g (96%) and micropore volume from 0.32 to 0.02cm³/g (94%). MEA-blocked AC particles adsorb selectively more CO₂ compared to non-impregnated AC particles by 172%, as the adsorption capacity increased from 49 to 18 mg/g, respectively.

b) Adsorption Kinetics
   i. Adsorption Rate
      Pseudo first and second order models were investigated to find out which model is predicting the adsorption rate appropriately.
   a) Pseudo First Order Model (SFO)
      Lagergren [22] published his SFO model to describe homogenous adsorption on solid phase as in equation 2. The equation depends on the adsorption capacity of an adsorption bed rather than concentration of adsorbate as in the case of first order model equation and the adsorption rate is related to the availability of adsorption sites [23]. It had been reported that SFO model can be applied adequately for the adsorption kinetics of CO₂ on AC [24].
The linear form of equation 1 is as in equation 3:

$$ln(q_e - q_t) = ln q_e - \frac{k_1}{2.303} t$$  

(3)

Where,

- $q_e$: Adsorption capacity at equilibrium, mg/g.
- $q_t$: Adsorption capacity at any time $t$, mg/g.
- $k_1$: Pseudo-first order rate constant, 1/min.
- $t$: Time, min.

Figure 1 is a plot of $ln (q_e - q_t)$ against $t$ for the whole adsorption experiment showing that the straight line fitting the experimental results has good value of $R^2$ equal to 0.9967, which is suggesting that the SFO kinetic model is applicable for this research. The slope of the straight line from equation 3 is $(\frac{k_1}{2.303})$, where the value of the SFO rate ($k_1$) was found to be 0.00552721/min.

**Figure 1**: Plot of $ln (q_e - q_t)$ against time

![Figure 1](image-url)

b. Pseudo Second Order Model (PSO)

PSO model is usually applied for chemisorption kinetics sorption of liquid solutions [25], the model is in equation 4 and its linear form is as in equation 5:

$$\frac{dq_t}{dt} = k(q_e - q_t)^2$$  

(4)

$$\frac{t}{q} = \frac{1}{k^2 q_e^2} + \frac{1}{q_e} t$$  

(5)

The plot of $t/q$ against $t$ for PSO model as in equation 5, didn’t yield straight line for the whole experimental data or even for any of its portions on contrary to the straight line of plot $ln (q_e - q_t)$ against $t$ for PFO model, which covers the whole experimental data and suggesting that the adsorption rate here is following PFO model which would be applied to calculate adsorption rate constant ($k_1$).

ii. Verification the Rate-Determining Step of the MEA-Impregnated AC Adsorption Bed

Equation 6 is the intraparticle diffusion equation [26].

$$q_t = k_p t^{\frac{1}{2}} + c$$  

(6)

Where:

- $q_t$: Amount of adsorbate absorbed at any time, mg g$^{-1}$.
- $k_p$: Intraparticle diffusion rate constant, mg g$^{-1}$ min$^{-1/2}$.

The multilinearity displayed in Figure 2, the plot of the amount of CO$_2$ adsorbed ($q_t$) against the square root of time ($t^{1/2}$) is suggesting that more than one step is taking place. The straight line fitting the curve is not passing through the origin, indicating that the intraparticle diffusion is not the adsorption rate restrictive step [27]. The curve is divided into three zones where the slope of the linear part indicating the rate of adsorption and the rate controlling step is represented by the linear section with lowest slope value [28]. The first zone is the initial zone where the external diffusion of CO$_2$ molecules through the bulk gas phase is taking place and the slope which is representing the adsorption rate is low. The second zone is the film diffusion where the mass transfer of CO$_2$ molecules is continuing through the CO$_2$ film surrounding the AC particles. The slope of the straight line of the second zone is higher than that of the initial zone but lower than that of the third zone due to the resistance exerted by CO$_2$ gas film, which is indicating that this step combining with the initial zone step are slow and the overall adsorption rate is controlled by mass transfer and film resistances respectively. In the third zone, where the amount of CO$_2$ molecules adsorbed ($q_t$)
versus $t^{1/2}$ is displaying a straight line with high slope value indicating that CO$_2$ molecules intraparticle diffusion step is fast, where CO$_2$ molecules adsorption is enhanced by the fast CO$_2$-MEA reaction. The intercept $(c)$ is an indicator of the thickness of the boundary layer surrounding the MEA-impregnated AC particles. Higher values of intercept suggest that the boundary layer is building up as the value of the $c$ in the initial zone is less than that of zone 1, which is in turn less than that of zone 2 deducing that diffusion through the gas film may be considered as the controlling step [29]. CO$_2$ adsorption in zone 2 is approaching its final stage and the active sites on MEA-impregnated AC particles are not able to adsorb more CO$_2$ molecules.

iii. **Gas Film Diffusion Model**

The transportation of CO$_2$ molecules from the gas stream bulk to the surface of the AC particles is playing a major role as the analysis of the intraparticle model showed that the mass transfer of CO$_2$ molecules through the gas film is the limiting step of CO$_2$ adsorption. To further inspect that gas film is the limiting step in CO$_2$ molecules adsorption, gas film diffusion model was applied [30], [31] and [32]:

$$\frac{q_t}{q_e} = 1 - e^{k_{fd} t}$$  \hspace{1cm} (7)

The linearized form of equation 7 is as in equation 8:

$$\ln(1 - F) = - k_{fd} t$$  \hspace{1cm} (8)

Where,

- $F$: Fractional adsorption equilibrium ($F = \frac{q_t}{q_e}$).
- $k_{fd}$: Film diffusion coefficient, min$^{-1}$.

A plot as in Figure 3 of $-\ln (1 - F)$ vs $t$ with intercept equal to zero and $R^2$ equal to 0.99 is suggesting that adsorption kinetics is controlled by diffusion through the CO$_2$ gas film surrounding the AC particles.

**Figure 2:** CO$_2$ Amount adsorbed, mg/g against square root of time, min$^{1/2}$

**Figure 3:** Plot of $-\ln (F-1)$ against time
iv. **Avrami (JMAK) Model**

Johnson-Mehl-Avrami-Kolmogorov (JMAK) model, which is called Avrami model too, is expressed in equation 9, [33] and [34]. Avrami equation describes the growth of crystallites with respect to time. In this work Avrami equation is describing the increasing numbers of CO₂ molecules by adsorption inside the AC pores.

\[
\alpha = 1 - \exp (-k_{Av} (t)^n)
\]  

(9)

Where, \( \alpha \) is adsorption fraction at time \( t \), \( k_{Av} \) is the Avrami kinetic constant, and \( n \) is a constant which represents the mechanism of particles adsorption (growth).

The linearized form is as in equation 10:

\[
\ln(-\ln(1-\alpha)) = n\ln k_{Av} + n\ln t
\]

(10)

### Figure 4: Plot of ln (-ln (1-\( \alpha \))) against ln (t)

Plotting ln (-ln (1-\( \alpha \))) against ln (t) as in Figure 4 producing straight line (\( R^2=0.9991 \)) with intercept equal to \( n\ln k_{Av} \) and slope equal to \( n \). If Avrami constant \( n \) equal to 1. Furthermore the value of Avrami exponent \( n \), which is \( 1 \leq n \leq 2 \) suggesting one dimensional growth of crystallites and that the growth is homogenous [35], which is agreeing with exponent \( n \) in micropore filling method of Dubinin-Astakhov (D-A), equation 11 and its linearized form equation 13. D-A equation is applicable for homogeneous carbonaceous adsorbents with micropore structures [36]. It was found in other study [37] that the value of D-A exponent \( n \) for MEA-impregnated beds is showing less heterogeneity and more homogeneity with their exponent \( n \) value equal to 2, where the value of exponent \( n \) in AC is 3 - 1.5. Moving from 3 to 1.5 the microporous system would be getting more heterogeneous [38] and [39].

---

**c) Mass Balance Mathematical Modeling**

Adsorption of CO₂ molecules from feed gas stream containing 15% CO₂ and 85% N₂ was performed in fixed bed packed column of non-impregnated and MEA-impregnated AC beds. Breakthrough time was employed as real time tool to evaluate the efficiency of the adsorption beds. CO₂ monitor was used to display the concentration (%) of the gas stream exiting the adsorption column. Graphs of CO₂ molecules concentration leaving the adsorption column plotted against time were obtained from the data acquisition logger connected to the outlet of the adsorption column.

i. **Mathematical Modeling of MEA-Impregnated 500 \( \mu \)m Adsorption Bed**

To formulate a general mathematical model corresponding to the mainly micropore adsorption mechanism and to cover the two stages mentioned earlier, the following assumptions were made:

1. The system operates under isothermal, isobaric and diabatic conditions.
2. The porosity of the adsorption bed was uniform and constant.
3. The equilibrium of adsorption is a nonlinear isotherm.
4. The velocity distribution is constant across the column diameter.
5. The volumetric flow rate is constant along the column.

Summarizes of the experimental parameters and simulation boundary conditions for the mathematical model validation are in Table 1:
Table 1: Experimental data and simulation boundary conditions

<table>
<thead>
<tr>
<th>Operating conditions</th>
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<td>Pressure</td>
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<tr>
<td>Temperature</td>
<td>25°C</td>
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<tr>
<td>Inlet concentration</td>
<td>6.05118E10⁻⁶ mol/ml</td>
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<tr>
<td>Inlet volumetric flow rate</td>
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<table>
<thead>
<tr>
<th>Adsorption column</th>
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<tbody>
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<td>Material</td>
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</tr>
<tr>
<td>Inside diameter</td>
<td>1 cm</td>
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<tr>
<td>Bed height</td>
<td>9 cm</td>
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<tr>
<td>Bed weight</td>
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<tr>
<td>Bed Volume</td>
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<table>
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<tr>
<th>Adsorbent properties</th>
<th>Non-impregnated AC</th>
<th>MEA-impregnated AC</th>
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<tbody>
<tr>
<td>Bed type</td>
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<td></td>
</tr>
<tr>
<td>Particles size</td>
<td>500 µm</td>
<td>500 µm</td>
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<tr>
<td>Micropore surface area</td>
<td>675 m²/g</td>
<td>65 m²/g</td>
</tr>
<tr>
<td>Micropore particle porosity</td>
<td>0.0956 m²/g</td>
<td>0.020 m²/g</td>
</tr>
<tr>
<td>Porosity</td>
<td>0.684</td>
<td>0.620</td>
</tr>
<tr>
<td>Bulk density</td>
<td>1.6387 cm³/g</td>
<td>1.6228 cm³/g</td>
</tr>
<tr>
<td>Bed Volume</td>
<td>7.2285 cm³</td>
<td>6.2857 cm³</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pseudo 1st order reaction constant (k)</th>
<th></th>
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<tbody>
<tr>
<td></td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>0.004836 1/min</td>
</tr>
</tbody>
</table>

ii. Mathematical Model of MEA-Impregnated AC Bed

The mathematical model was based on the CO₂ molecules breaking through the adsorption bed. The adsorption of CO₂ molecules was declining and more CO₂ molecules were exiting the bed.

The general equation of mass balance with first order chemical reaction for CO₂ in the feed gas:

Accumulation = Input – Output + Generation

As the mass balance would be conducted on CO₂ molecules exiting the adsorption bed, the mass balance equation would be:

Output = Input - Accumulation + Generation

\[
v \frac{dc_v}{dt} = qc_i - qc_{acc} + vk_1c
\]  

(14)

Where:
- \(K_1\): Pseudo first order reaction constant, 1/min.
- \(C_i\): Concentration of CO₂ entering the AC bed, mol/ml.
- \(C_v\): Concentration of CO₂ exiting the AC bed, mol/ml.
- \(C_{acc}\): Concentration of CO₂ accumulated in the AC bed, mol/ml.

Equation 14 is an ordinary first order linear differential equation and the final solution would be as in Equation 15.

\[
C = C_i + n e^{k_1 t} = C_i + n e^{(-q/v - k_1) t} 
\]

(15)

Initial boundary condition:
At \(t = 0\), \(C = 0\)
\(n = -C_i\)

Then equation 15 would be:

\[
C = C_i - C_i e^{(-q/v - k_1) t} 
\]

(16)

Rearranging equation 16:

\[
C = C_i \left(\frac{q/v}{(q/v) - k_1}\right) \left(1 - e^{-(q/v - k_1) t}\right) 
\]

(17)

The simulated results were validated by using the experimental results of the MEA-impregnated activated carbon bed. The simulated results were compared with the experimental data. The simulated data demonstrated a reasonable agreement with the experimental data, as the root mean square error (RMSE) calculated was 6.75915E-06. The simulated and experimental data of MEA-impregnated AC beds were plotted in Figure 5.
IV. Conclusions

MEA-impregnated AC particles were used to adsorb \( \text{CO}_2 \) from gas mixture. Results are showing that AC particles impregnated with MEA adsorb \( \text{CO}_2 \) molecules in a pseudo first order reaction manner and that the controlling step in this reaction is the mass transfer of \( \text{CO}_2 \) molecules from through the \( \text{CO}_2 \) gas film and not the intraparticle diffusion of \( \text{CO}_2 \) molecules inside the pores of MEA-impregnated AC particles. Due to the homogeneity of the MEA-impregnated activated carbon particles the adsorption of \( \text{CO}_2 \) molecules follows the Avrami model of homogenous crystallites growth. The mass balance mathematical model showed that the experimental and simulated breakthrough curves have good agreement, as the root mean square error (RMSE) was 5.7678E-09 and 3.88532E-09 for non-impregnated and MEA-impregnated beds respectively, which also proved that the adsorption mechanism of both beds is the same.

Acknowledgment

The authors would like to thank University of Malay for offering the necessary fund for this research through the, University of Malaya Research grant UMRG RP15/2012A.

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Moisture-dependent Physical Properties and Hydration Kinetics of Peanut Kernel

By Shubhra Shekhar, Manoj Singh & Kamlesh Prasad

Abstract- The physical properties of agricultural objects affected by moisture are essential for effective postharvest unit operations. Raw peanut in India is used in a wide variety of forms, and the process involved often soaking in water before being used as an ingredient in the preparation of different delicacies. Soaking and grinding in preparation of seasonings are commonly used to garnish the most traditional fermented foods such as idly, dosa, vada, and uttapam with several other breakfast items in the Indian subcontinent. The effect of moisture content (6.57 to 35.07%) on physical properties and the temperature (5 to 35°C) dependent soaking behavior of peanut kernel of GG-20 genotype were assessed. The increase in moisture content of peanuts has affected the dimensional characteristics linearly, whereas most of the gravimetric and frictional properties followed a nonlinear trend. The feasibility of different models describing the hydration behavior was assessed for the peanut kernel. The adequacy of fitted models was determined using the coefficient of determination ($R^2$), chi-square ($\chi^2$), and root mean square error (RMSE). The values of these parameters as higher $R^2$ ($\geq 0.96$) and lower $\chi^2$ ($\leq 0.005$) and RMSE ($\leq 0.020$) reflect the applicability of models in describing the soaking behavior of the peanut kernel.

Keywords: peanut, physical properties, soaking, diffusion, models, hydration.

GJRE-C Classification: DDC Code: 551.571 LCC Code: QC915

Strictly as per the compliance and regulations of:
Moisture-dependent Physical Properties and Hydration Kinetics of Peanut Kernel

Shubhra Shekhar *, Manoj Singh ° & Kamlesh Prasad °

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Keywords: peanut, physical properties, soaking, diffusion, models, hydration.

I. INTRODUCTION

Peanut (Arachis hypogaea) is the 3rd most crucial oilseed crop after soybean and cotton. Being a legume crop, it is rich in protein. This tropical cash crop is often referred to as “The king of oilseeds,” groundnut, earth nut, wonder nut, or poor men’s cashew nut. India is the pioneer producer of peanuts, with a production of 6.48 MMT and contributed more than 18% of world production for 2020 (FAOSTAT 2022)[11]. The chief peanut-producing states in India are Gujarat, Rajasthan, Andhra Pradesh, Tamil Nadu, and Madhya Pradesh. These five states account for around 90% of the total area under peanut cultivation, mainly as the Kharif season crop (June - September). As a leguminous crop, it is grown in crop rotation mainly to maintain soil health and reduce soil erosion.

Eighty percent of the total peanut produced in India undergoes processing for oil extraction through either mechanical or solvent extraction in a batch or continuous process. The high temperature in most efficient mechanical extractors damages the edible quality of obtained defatted materials and is unsuitable for human consumption (BADWAIK, PRASAD, DEKA 2012)[2]. Residual solvent associated with solvent-extracted defatted peanut meal creates problems in its utilization for various value-added products (BADWAIK, PRASAD, SETH 2014)[3], therefore affecting the loss of valuable resources.

Peanut may be considered a functional food, as it contains numerous functional and health-promoting compounds, including arginine, mono, and polyunsaturated fatty acids, dietary fiber, folate, niacin, vitamin E, and most of the vital minerals (GOPALAN, RAMA SASTRI, BALASUBRAMANIAN 1971)[6] beneficial for human health. The presence of bioactive phenolic compounds provides this valuable biomaterial with a rich source of antioxidants (FRANCISCO, RESURRECCION 2008)[5]. Raw peanut in India is used in a wide variety of forms. The process involved often soaking in water before being used as an ingredient in the preparation of different delicacies. Soaking and grinding in preparation of seasonings are commonly used to garnish the most traditional fermented foods such as idly, dosa, vada, and uttapam with several other breakfast items in the Indian subcontinent (ACHAYA 1984)[8].

The physical properties of peanut kernels, like those of other seeds, become helpful in designing equipment, especially for handling, processing, and storing. The moisture content of grain affects the efficiency of cleaning, grading, and separation processes (YALÇIN, ÖZARSLAN, AKBAŞ 2007)[7]. The author has made investigations the determination of the physical properties of various leguminous splits, such as chickpeas, lentils, black gram, and green gram (BHATIA, SHARMA, PRASAD, PRASAD 2009)[8]. The author has made investigations the determination of the physical properties of various leguminous splits, such as chickpeas, lentils, black gram, and green gram (BHATIA, SHARMA, PRASAD, PRASAD 2009)[8]. The high temperature in most efficient mechanical extractors damages the edible quality of obtained defatted materials and is unsuitable for human consumption (BADWAIK, PRASAD, DEKA 2012)[2]. Residual solvent associated with solvent-extracted defatted peanut meal creates problems in its utilization for various value-added products (BADWAIK, PRASAD, SETH 2014)[3], therefore affecting the loss of valuable resources.
Limited studies on water absorption kinetics for peanuts were carried out. The present study aims to investigate the moisture-dependent physical properties of temperature-dependent water absorption kinetics to assess the adequacy of mathematical models in describing the process of water absorption kinetics.

II. MATERIALS AND METHODS

a) Sample Preparation

The dried whole peanut pods of genotype GG-20 were procured from the Central Institute of Postharvest Engineering and Technology (CIPHET), Punjab Agricultural University Campus (PAU), Ludhiana (Punjab). The whole peanut pods were cleaned and dehulled. The kernels were manually sorted to separate damaged and shrunk kernels. The sorted sound kernels (Fig. 1) were stored in air-tight glass containers and under refrigerated conditions. The samples were taken out as per the experimental requirements. The initial moisture contents of the kernels were determined using the hot air oven method (GUPTA, DAS 2000) [14]. The predetermined weight of moisture was impregnated to obtain the four levels of moisture content for the peanut kernels as 5.57±0.56%, 15.09±0.65%, 24.98±0.51%, and 35.07±0.61% (wet basis), conditioned and stored at 5±1°C. The samples for the study were taken out and kept in the ambient environmental condition in sealed pouches to equilibrate quickly and without any change of moisture content.

b) Determination of Physical Properties

The geometrical shape consisting of a cylinder and two hemispheres at the ends has been found to be quantitative appropriateness for peanuts (AKCALI, INCE, GUZEL 2006) [19]. The physical dimensions were determined randomly measured for three major perpendicular dimensions, length (L), width (W), and thickness (T), of 100 seeds of each variety using a digital vernier caliper (Mitutoyo Corporation, Japan) having the precision of 0.01mm. The geometric mean dimension (Dg) of peanut kernels was found using the relationship as given (MOHSENIN 2020) [16]:

\[ D_g = \sqrt[3]{LWT} \]  

(1)

The criteria used to describe the shape of the seed are sphericity. Thus, the sphericity (ϕ) was accordingly computed (MOHSENIN 2020) [16] as:

\[ \phi = \frac{D_g}{L} \times 100 \]  

(2)

The surface area (Sa) was calculated (Eqn. 3) considering the geometrical properties of the kernel as:

\[ S_a = \pi \times L \times T \]  

(3)

Electronic balance (Ishida Co. Ltd., Japan) was used to measure the weight of the sample to an accuracy of 0.001 g. The true density of a peanut kernel is defined as the ratio of the mass of the seed to the solid volume occupied (DESHPANDE, BAL, OJHA 1993) [17]. The liquid displacement technique determined the seed volume and its true density. The immersion time was maintained below 10 sec, which was considered too small to absorb water. The amount of displaced water was recorded from the graduated scale of the measuring cylinder (AMIN, HOSSAIN, ROY 2004) [18]. The porosity (ε) of bulk seed was computed from the values of true density (ρt) and bulk density (ρb) using the relationship (Eqn. 4) given (MOHSENIN 2020) [16]:

\[ \epsilon = \left[ 1 - \left( \frac{\rho_b}{\rho_t} \right) \right] \times 100 \]  

(4)

To determine the angle of repose (ϕ), a cylinder of 50 mm diameter was used. The height and diameter of the formed heap of peanut kernels were measured to determine the angle of repose (ϕ) using the relationship:

\[ \varphi = \arctan \left( \frac{2H}{D} \right) \]  

(5)

Where H is the height of the heap (mm), and D is the diameter of the heap (mm) at its base.

The static coefficient of friction (μ) was determined for three structural surfaces, namely galvanized steel sheets, glass, and plywood. A plastic cylinder of 50 mm diameter and 60 mm height was placed on an adjustable tilting flat plate faced with the test surface and filled with a sample of about 100 g. The cylinder was raised slightly to ensure complete contact with the peanut kernel and the experimenting surface and to avoid any error being induced by the test cylinder. The setup was then inclined gradually until the cylinder started sliding down. The angle of tilt was noted from a graduated scale (DUTTA, NEMA, BHARDWAJ 1988) [19]. SHERPHED, BHARDWAJ 1986) [20].

\[ \tan \alpha = \mu \]  

(6)

All the experiments were replicated at least ten times, and average values were reported.

c) Water Absorption during Soaking

The hydration behavior of peanut kernel was determined by soaking a 50g sample in cylindrical glass containers containing 250 ml of distilled water (1:5 w/v) (ABD EL-HADY, HABIBA 2003) [21]. Soaking water temperatures were 5, 15, 25, and 35°C. Before each experiment, the sample, containers, and distilled water were kept at the desired temperature for a few hours. Hydration behavior was precisely observed for up to 24 hours using an electronically controlled water bath to an accuracy of ±0.2°C (RESIO, AGUERRE, SUAREZ 2005) [22]. The moisture content of the sample during hydration was determined based on the sample's weight increase. Soaked kernels were immediately blotted with the muslin cloth to remove any chance of the presence...
of surface water before taking the reading (VERMA, PRASAD 1999)[23].

d) Modeling of Water Absorption of Peanut Kernels

The kinetics of hydration of peanut kernels was assessed using selected models presented in Table 1. Analysis of absorption behavior through selected models, the time-dependent data of moisture content, and moisture ratio (MR) of peanut kernels is essential. The moisture ratio was calculated as follows:

\[ MR = \frac{M_t - M_e}{M_0 - M_e} \]  \hspace{1cm} (7)

These parameters can be calculated as follows:

\[ R^2 = \frac{\sum_{i=1}^{N} (MR_{exp,i} - MR_{pre,i})^2}{\sum_{i=1}^{N} (MR_{pre,i})^2} \]  \hspace{1cm} (8)

\[ \chi^2 = \frac{1}{N-n} \sum_{i=1}^{N} (MR_{exp,i} - MR_{pre,i})^2 \]  \hspace{1cm} (9)

\[ \text{RMSE} = \frac{1}{N} \sqrt{\sum_{i=1}^{N} (MR_{exp,i} - MR_{pre,i})^2} \]  \hspace{1cm} (10)

where \( MR_{exp,i} \) is the \( i^{th} \) experimentally observed moisture ratio, \( MR_{pre,i} \) the \( i^{th} \) predicted moisture ratio, \( N \) is the number of observations, and \( n \) is the number of constants in the model (YALÇın, ÖZARSLAN, AKBAŞ 2007)[27], TOĞRUL, PEHLIVAN 2003)[28].

III. Results and Discussion

a) Physical Properties

An increase in moisture content from 5 to 35% has shown a linear change in length (L), width (W), thickness (T), and geometric mean diameter (GMD) of peanut kernels (Fig. 2) from 14.52 to 16.37 mm, 8.40 to 9.61 mm, 7.41 to 8.79 mm, and 9.67 to 11.14 mm, respectively (Table 2). A linear dimensional increase in peanut kernels with an increase in moisture content was mainly due to the absorption of moisture that resulted in the swelling of peanut cells and thus affected kernel's dimensional properties. The linear increase in width and thickness has also been reported for popcorn kernel (KARABABA 2006)[29] and pistachio nut (KASHANINEJAD, MORTAZAVI, SAFEKORDI, TABIL 2006)[27]. A minor initial decrease in the sphericity (\( \Phi \)) of peanut kernel with further increase as the increase in moisture content is evident (Fig 2). The initial decrease in sphericity may be attributed to the initial higher rate of lateral expansion in comparison to the change in the width and thickness. However, at later stages of moisture gain, the rate of expansion in width and thickness increased more and caused to increase in sphericity. The high sphericity value suggests that the kernel tends towards a spherical shape (OMOBUWAOJO, SANNI, OLAJIDE 2000)[28], being cylindrical. The surface area (\( S_a \)) of peanut kernels increased linearly from 3.38 to 4.52 cm² with increased moisture content. It was of overall 33.73% increase (Fig. 2). A surface area is a relevant tool in determining the shape of the kernel and indicating the material behavior on oscillating surfaces during processing (ALONGE, ADIGUN 1999)[29].

The unit mass (UM) of peanut kernel was found to be increased from 0.462 to 0.724 gm with an increase in moisture content (Table 2). Differential sphericity, surface area, and unit mass of materials serve in designing aerodynamic cleaning systems. Bulk density (\( \rho_b \)) decreased from 560 to 496 kg/m³ with an increase in moisture (Fig. 3). This decrease in the bulk density on change of moisture content is probably due to higher volume expansion in kernel in comparison to weight change on change in moisture content (Fig. 3). A decrease in bulk density was found for chickpea seeds from 600 to 741.4 kg/m³ with an increase in moisture content from 4.9 to 14.1% (AMIN, HOSSAIN, Roy 2004)[18]. A similar relation between moisture content and bulk density was reported for Soybean (DESHPANDE, BAL, OJHA 1993)[17], sunflower seed (GUPTA, DAS 1997)[30], and popcorn kernel (KARABABA 2006)[29]. Porosity increased linearly from 39.58 to 46.77% with an increase in moisture content, and similar trends were reported for gram (DUTTA, NEMA, BHARDWAJ 1988)[31], sunflower kernel (GUPTA, DAS 1997)[30], green gram (NIMKAR, CHATTOPADHYAY 2001)[32], chickpea seed (KONAK, CARMAN, AYDIN 2002)[33], and popcorn kernel(KARABABA 2006)[29].

The angle of repose and coefficient of friction as frictional properties are essential in designing equipment for solid flow and storage structures. The coefficient of friction between the seed and wall is an essential...
The angle of repose of peanut kernels increased from 39.01 to 50.48° with an increase in moisture content in the range of 5 to 35% (Table 2). The static coefficient of friction at various moisture levels reflects the increase in moisture content for all the analyzed surfaces (Fig. 4). This is due to the increased adhesion properties between the seed and the material surfaces at higher moisture levels.

**b) Soaking Kinetics**

The behavior of moisture uptake during time-dependent soaking by peanut kernel at temperatures 5, 15, 25, and 35°C is shown in Fig. 5. An early rapid hydration rate in the primary phase with a slower hydration rate in the second phase was observed (Fig. 5). The time needed for the transition of phase was found by applying the creep deformation properties in the water absorption process, observed to be temperature-dependent, and were 2.80, 2.56, 1.89, and 1.24 hrs at 5, 15, 25, and 35°C, respectively. The diffusion process governs the rapid initial water uptake in the primary phase was probably due to the filling of capillaries on the surface of the seed coats. A decrease in water absorption rate further in the relaxation phase may be attributed to the reason that water filling decreased the driving force. At any time, the moisture content of the kernels increased with temperature, indicating that the amount of water absorbed during hydration was a function of both soaking time and temperature (Fig. 5). The probabilistic Weibull model describes a system's behavior with some degree of variability (CUNHA, OLIVEIRA, OLIVEIRA 1998)\[^{38}\]. Weibull shape parameter (α) is a behavior index that depends on the process mechanism, and the higher its value, the slower the process in the initial phase of absorption (CUNHA, OLIVEIRA, OLIVEIRA 1998)\[^{38}\]; MACHADO, OLIVEIRA, GEKAS, SINGH 1998\[^{39}\]. Thus, the inverse of the Weibull shape parameter was thus considered an indicator of the initial rate of the hydration process. An increase in the temperature of soaking thus reduced the value of α from 5.609 to 1.546, further justifying the obtained temperature-dependent soaking behavior for peanut kernel (Table 3).

Weibull scale parameter (β) represents the time needed to accomplish approximately 63% of total absorbed water (CUNNINGHAM, MCMINN, MAGEE, RICHARDSON 2007)\[^{40}\]. Table 3 shows the effect of temperature change on relative rates of moisture diffusion and thus affecting the overall absorption process. This implies that with an increase in temperature from 5 to 35°C, the absorption rate in the primary phase of soaking was accelerated significantly (Fig. 5).

Fitting the exponential model to the experimental data at four different temperatures for the hydration process (Fig. 6) depicts an increasing trend of hydration rate constant (K) with the soaking temperature (Table 3). The high observed value of the coefficient of determination (R²) confirmed the applicability in the prediction of the soaking process of the peanut kernel.

A large amount of absorbed water is present in the first phase of the process and further reaches equilibrium in the relaxation phase of soaking. However, at lower temperatures, the absorbed water in the primary soaking phase is less, and the absorption continues in the second phase. The values for the parameters K₁, K₂; K, and α & β in Peleg, Exponential, and Weibull models, respectively, supported the findings.
ii. Diffusivity Model and Activation Energy

To predict moisture diffusivity during the soaking of peanut kernels, the second Fick’s law solution for diffusion out of an infinite cylinder being length of more than twice the width was used. For this purpose, the assumptions considered were (i) the effective diffusion coefficient is independent of moisture concentration, (ii) the volume of the kernel does not change during water absorption, and (iii) the surface of the kernel reaches the equilibrium moisture content instantaneously upon immersion in absorption media.

The general series solution of Fick’s second law in the infinite cylinder is given below (CRANK 1975)[41]:

\[
MR = \sum_{n=1}^{\infty} \frac{4}{\beta_n^2} \exp \left[ -\frac{\beta_n^2}{R^2} \times D_e t \right]
\]

(11)

\[
D_e = D_0 \exp \left( -\frac{E_a}{RT_a} \right)
\]

(12)

\[
D_0 \text{ is the effective diffusivity (m}^2\text{/sec.) and } R \text{ is the characteristic radius of the cylinder. Eqn calculated the effective diffusion coefficient. (11) using slopes derived from the linear regression of ln (MR) against time data. The effective diffusion coefficient during the soaking of peanut kernels varied from 2.9053×10-10 to 10.753×10-10 (m2/sec.) over the temperature range studied 5-35°C (Table 4).

A similar result was also obtained with other agricultural seeds. The effectiveness of diffusivity during soaking of rice varied from 5.58×10-11 to 3.57×10-11 (m2/sec.) in the temperature range from 25 to 70°C (KASHANINEJAD, MAGHSOUDLOU, RAFIGE, KHOMEIRI 2007)[43], in case of soybean, varied from 1.08×10-10 to 2.00×10-10 m2/sec. in the temperature range from 40 to 60°C (HSU 1983)[42], and 1.632×10-9 to 3.237×10-9 (m2/sec.) in case of splits chickpea at the hydration temperature range of 40 to 60°C (PRASAD, VAIRAGAR, BERA 2010)[9].

The effect of temperature on effective diffusivity is generally described using the Arrhenius-type relationship to obtain a better agreement of the predicted curve with experimental data:

\[
D_e = D_0 \exp \left( -\frac{E_a}{RT_a} \right)
\]

(12)

The hydration kinetics of peanut kernel (GG-20) in plain water at different temperatures was examined. The soaking process exhibited an initial faster rate of water absorption followed by a slower pace during the secondary absorption phase. The transition from primary absorption phases occurred approximately between 1.23 to 2.80 hrs, depending on the soak water temperature, which may be used as the deciding time for soaking of peanut. The moisture absorption was an endothermic process with heat absorption of 32.175 KJ/mol.

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**Fig. 1:** Charged coupled device snapshot showing shelled peanut kernels of GG-20 variety

**Fig. 2:** Moisture dependent dimensional characteristics of peanut kernels


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**Fig. 3:** Moisture dependent gravimetric characteristics of peanut kernels

**Fig. 4:** Moisture dependent frictional characteristics of peanut kernels
Fig. 5: Temperature dependent hydration characteristics of peanut kernel (Peleg and Weibull fit)

Fig. 6: Temperature dependent hydration characteristics of peanut kernel (Exponential fit)
Reciprocal of absolute temperature, $T_a^{-1}, K^{-1}$

![Fig. 7: Arrhenius plot of effective diffusivity of peanut kernels](image)

Table 1: Models used for describing the peanut kernel hydration data

<table>
<thead>
<tr>
<th>Model Name</th>
<th>Equation</th>
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<tr>
<td>Peleg</td>
<td>$M = M_0 + \frac{t}{(K_1 + K_2 t)}$</td>
</tr>
<tr>
<td>Exponential</td>
<td>$MR = \exp(-kt)$</td>
</tr>
<tr>
<td>Weibull</td>
<td>$MR = \exp(-\frac{t}{\lambda})^\beta$</td>
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</tbody>
</table>

Table 2: Moisture dependent physical properties of peanut kernel (GG-20)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Moisture Levels of shelled peanut (% wwb)</th>
<th>6.57±0.44%</th>
<th>15.74±0.01%</th>
<th>25.21±0.03%</th>
<th>35.07±0.09%</th>
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</thead>
<tbody>
<tr>
<td>Length, mm</td>
<td></td>
<td>14.52±1.48</td>
<td>15.74±1.41</td>
<td>15.78±1.52</td>
<td>16.37±1.20</td>
</tr>
<tr>
<td>Width, mm</td>
<td></td>
<td>8.40±0.72</td>
<td>8.61±0.44</td>
<td>9.22±0.49</td>
<td>9.61±0.47</td>
</tr>
<tr>
<td>Thickness, mm</td>
<td></td>
<td>7.41±0.43</td>
<td>7.76±0.58</td>
<td>7.92±0.65</td>
<td>8.79±0.70</td>
</tr>
<tr>
<td>GMD, mm</td>
<td></td>
<td>9.67±0.66</td>
<td>10.17±0.56</td>
<td>10.48±0.54</td>
<td>11.14±0.50</td>
</tr>
<tr>
<td>Sphericity, %</td>
<td></td>
<td>66.60±4.20</td>
<td>64.61±3.66</td>
<td>66.44±4.97</td>
<td>68.03±4.56</td>
</tr>
<tr>
<td>Surface Area, cm²</td>
<td></td>
<td>3.38±0.45</td>
<td>3.84±0.50</td>
<td>3.93±0.47</td>
<td>4.52±0.43</td>
</tr>
<tr>
<td>Unit Mass, gm</td>
<td></td>
<td>0.462±0.161</td>
<td>0.565±0.127</td>
<td>0.567±0.139</td>
<td>0.724±0.143</td>
</tr>
<tr>
<td>Bulk Density, kg/m³</td>
<td></td>
<td>560.0±10</td>
<td>491.0±7.0</td>
<td>460.0±12.0</td>
<td>496.0±17.0</td>
</tr>
<tr>
<td>True Density, gm/ml</td>
<td></td>
<td>0.927±0.010</td>
<td>0.877±0.016</td>
<td>0.865±0.019</td>
<td>0.932±0.026</td>
</tr>
<tr>
<td>Porosity, %</td>
<td></td>
<td>39.58±0.95</td>
<td>43.98±1.63</td>
<td>46.81±1.09</td>
<td>46.77±0.97</td>
</tr>
<tr>
<td>Angle of Repose, degree</td>
<td></td>
<td>39.01±1.87</td>
<td>49.35±1.55</td>
<td>50.09±1.06</td>
<td>50.48±0.68</td>
</tr>
<tr>
<td>CSF (Steel)</td>
<td></td>
<td>0.248±0.004</td>
<td>0.331±0.014</td>
<td>0.367±0.008</td>
<td>0.406±0.006</td>
</tr>
<tr>
<td>CSF (Glass)</td>
<td></td>
<td>0.158±0.007</td>
<td>0.337±0.020</td>
<td>0.379±0.010</td>
<td>0.409±0.006</td>
</tr>
<tr>
<td>CSF (Plywood)</td>
<td></td>
<td>0.295±0.004</td>
<td>0.300±0.008</td>
<td>0.320±0.013</td>
<td>0.430±0.028</td>
</tr>
</tbody>
</table>
### Table 3: Estimation of the parameters and goodness of fit for Peleg, Exponential and Weibull models

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Peleg Model</th>
<th>Exponential Model</th>
<th>Weibull Model</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$K_1$</td>
<td>$K_2$</td>
<td>$R^2$</td>
</tr>
<tr>
<td>5</td>
<td>0.0</td>
<td>0.0</td>
<td>0.98</td>
</tr>
<tr>
<td>15</td>
<td>0.0</td>
<td>0.0</td>
<td>0.99</td>
</tr>
<tr>
<td>25</td>
<td>0.0</td>
<td>0.0</td>
<td>0.98</td>
</tr>
<tr>
<td>35</td>
<td>0.0</td>
<td>0.0</td>
<td>0.99</td>
</tr>
</tbody>
</table>

### Table 4: Temperature dependent effective diffusion coefficient of peanut kernel

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Effective diffusivity (m²/sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>$2.905 \times 10^{-10}$</td>
</tr>
<tr>
<td>15</td>
<td>$4.833 \times 10^{-10}$</td>
</tr>
<tr>
<td>25</td>
<td>$8.581 \times 10^{-10}$</td>
</tr>
<tr>
<td>35</td>
<td>$10.753 \times 10^{-10}$</td>
</tr>
</tbody>
</table>
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FELLOWS/ASSOCIATES OF ENGINEERING RESEARCH COUNCIL
FERC/AERC MEMBERSHIPS

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<table>
<thead>
<tr>
<th><strong>ASSOCIATE</strong></th>
<th><strong>FELLOW</strong></th>
<th><strong>RESEARCH GROUP</strong></th>
<th><strong>BASIC</strong></th>
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<td>$4800 lifetime designation</td>
<td>$6800 lifetime designation</td>
<td>$12500.00 organizational</td>
<td>APC per article</td>
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<td>Certificate, LoR and Momento Unlimited discounted publishing/year</td>
<td>Certificates, LoRs and Momentos Unlimited free publishing/year</td>
<td>GJ Community Access</td>
</tr>
<tr>
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<td>Gradation of Research Unlimited research contacts/day</td>
<td>Gradation of Research Unlimited research contacts/day</td>
<td>GJ Community Access</td>
</tr>
<tr>
<td>1 GB Cloud Storage</td>
<td>5 GB Cloud Storage</td>
<td>Unlimited Cloud Storage</td>
<td>GJ Community Access</td>
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</table>
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- First character must be three lines drop-capped.
- The paragraph before spacing of 1 pt and after of 0 pt.
- Line spacing of 1 pt.
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11. **Pick a good study spot**: Always try to pick a spot for your research which is quiet. Not every spot is good for studying.

12. **Know what you know**: Always try to know what you know by making objectives, otherwise you will be confused and unable to achieve your target.

13. **Use good grammar**: Always use good grammar and words that will have a positive impact on the evaluator; use of good vocabulary does not mean using tough words which the evaluator has to find in a dictionary. Do not fragment sentences. Eliminate one-word sentences. Do not ever use a big word when a smaller one would suffice.

Verbs have to be in agreement with their subjects. In a research paper, do not start sentences with conjunctions or finish them with prepositions. When writing formally, it is advisable to never split an infinitive because someone will (wrongly) complain. Avoid clichés like a disease. Always shun irritating alliteration. Use language which is simple and straightforward. Put together a neat summary.

14. **Arrangement of information**: Each section of the main body should start with an opening sentence, and there should be a changeover at the end of the section. Give only valid and powerful arguments for your topic. You may also maintain your arguments with records.

15. **Never start at the last minute**: Always allow enough time for research work. Leaving everything to the last minute will degrade your paper and spoil your work.

16. **Multitasking in research is not good**: Doing several things at the same time is a bad habit in the case of research activity. Research is an area where everything has a particular time slot. Divide your research work into parts, and do a particular part in a particular time slot.

17. **Never copy others’ work**: Never copy others’ work and give it your name because if the evaluator has seen it anywhere, you will be in trouble. Take proper rest and food: No matter how many hours you spend on your research activity, if you are not taking care of your health, then all your efforts will have been in vain. For quality research, take proper rest and food.

18. **Go to seminars**: Attend seminars if the topic is relevant to your research area. Utilize all your resources.

19. **Refresh your mind after intervals**: Try to give your mind a rest by listening to soft music or sleeping in intervals. This will also improve your memory. Acquire colleagues: Always try to acquire colleagues. No matter how sharp you are, if you acquire colleagues, they can give you ideas which will be helpful to your research.

20. **Think technically**: Always think technically. If anything happens, search for its reasons, benefits, and demerits. Think and then print: When you go to print your paper, check that tables are not split, headings are not detached from their descriptions, and page sequence is maintained.

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21. Adding unnecessary information: Do not add unnecessary information like "I have used MS Excel to draw graphs." Irrelevant and inappropriate material is superfluous. Foreign terminology and phrases are not apropos. One should never take a broad view. Analogy is like feathers on a snake. Use words properly, regardless of how others use them. Remove quotations. Puns are for kids, not grunt readers. Never oversimplify: When adding material to your research paper, never go for oversimplification; this will definitely irritate the evaluator. Be specific. Never use rhythmic redundancies. Contractions shouldn’t be used in a research paper. Comparisons are as terrible as clichés. Give up ampersands, abbreviations, and so on. Remove commas that are not necessary. Parenthetical words should be between brackets or commas. Understatement is always the best way to put forward earth-shaking thoughts. Give a detailed literary review.

22. Report concluded results: Use concluded results. From raw data, filter the results, and then conclude your studies based on measurements and observations taken. An appropriate number of decimal places should be used. Parenthetical remarks are prohibited here. Proofread carefully at the final stage. At the end, give an outline to your arguments. Spot perspectives of further study of the subject. Justify your conclusion at the bottom sufficiently, which will probably include examples.

23. Upon conclusion: Once you have concluded your research, the next most important step is to present your findings. Presentation is extremely important as it is the definite medium though which your research is going to be in print for the rest of the crowd. Care should be taken to categorize your thoughts well and present them in a logical and neat manner. A good quality research paper format is essential because it serves to highlight your research paper and bring to light all necessary aspects of your research.

Informal Guidelines of Research Paper Writing

Key points to remember:

- Submit all work in its final form.
- Write your paper in the form which is presented in the guidelines using the template.
- Please note the criteria peer reviewers will use for grading the final paper.

Final points:

One purpose of organizing a research paper is to let people interpret your efforts selectively. The journal requires the following sections, submitted in the order listed, with each section starting on a new page:

The introduction: This will be compiled from reference matter and reflect the design processes or outline of basis that directed you to make a study. As you carry out the process of study, the method and process section will be constructed like that. The results segment will show related statistics in nearly sequential order and direct reviewers to similar intellectual paths throughout the data that you gathered to carry out your study.

The discussion section:

This will provide understanding of the data and projections as to the implications of the results. The use of good quality references throughout the paper will give the effort trustworthiness by representing an alertness to prior workings.

Writing a research paper is not an easy job, no matter how trouble-free the actual research or concept. Practice, excellent preparation, and controlled record-keeping are the only means to make straightforward progression.

General style:

Specific editorial column necessities for compliance of a manuscript will always take over from directions in these general guidelines.

To make a paper clear: Adhere to recommended page limits.

Mistakes to avoid:

- Insertion of a title at the foot of a page with subsequent text on the next page.
- Separating a table, chart, or figure—confine each to a single page.
- Submitting a manuscript with pages out of sequence.
- In every section of your document, use standard writing style, including articles ("a" and "the").
- Keep paying attention to the topic of the paper.
• Use paragraphs to split each significant point (excluding the abstract).
• Align the primary line of each section.
• Present your points in sound order.
• Use present tense to report well-accepted matters.
• Use past tense to describe specific results.
• Do not use familiar wording; don't address the reviewer directly. Don't use slang or superlatives.
• Avoid use of extra pictures—include only those figures essential to presenting results.

Title page:
Choose a revealing title. It should be short and include the name(s) and address(es) of all authors. It should not have acronyms or abbreviations or exceed two printed lines.

Abstract: This summary should be two hundred words or less. It should clearly and briefly explain the key findings reported in the manuscript and must have precise statistics. It should not have acronyms or abbreviations. It should be logical in itself. Do not cite references at this point.

An abstract is a brief, distinct paragraph summary of finished work or work in development. In a minute or less, a reviewer can be taught the foundation behind the study, common approaches to the problem, relevant results, and significant conclusions or new questions.

Write your summary when your paper is completed because how can you write the summary of anything which is not yet written? Wealth of terminology is very essential in abstract. Use comprehensive sentences, and do not sacrifice readability for brevity; you can maintain it succinctly by phrasing sentences so that they provide more than a lone rationale. The author can at this moment go straight to shortening the outcome. Sum up the study with the subsequent elements in any summary. Try to limit the initial two items to no more than one line each.

Reason for writing the article—theory, overall issue, purpose.
• Fundamental goal.
• To-the-point depiction of the research.
• Consequences, including definite statistics—if the consequences are quantitative in nature, account for this; results of any numerical analysis should be reported. Significant conclusions or questions that emerge from the research.

Approach:
• Single section and succinct.
• An outline of the job done is always written in past tense.
• Concentrate on shortening results—limit background information to a verdict or two.
• Exact spelling, clarity of sentences and phrases, and appropriate reporting of quantities (proper units, important statistics) are just as significant in an abstract as they are anywhere else.

Introduction:
The introduction should "introduce" the manuscript. The reviewer should be presented with sufficient background information to be capable of comprehending and calculating the purpose of your study without having to refer to other works. The basis for the study should be offered. Give the most important references, but avoid making a comprehensive appraisal of the topic. Describe the problem visibly. If the problem is not acknowledged in a logical, reasonable way, the reviewer will give no attention to your results. Speak in common terms about techniques used to explain the problem, if needed, but do not present any particulars about the protocols here.

The following approach can create a valuable beginning:
• Explain the value (significance) of the study.
• Defend the model—why did you employ this particular system or method? What is its compensation? Remark upon its appropriateness from an abstract point of view as well as pointing out sensible reasons for using it.
• Present a justification. State your particular theory(-ies) or aim(s), and describe the logic that led you to choose them.
• Briefly explain the study's tentative purpose and how it meets the declared objectives.
Approach:

Use past tense except for when referring to recognized facts. After all, the manuscript will be submitted after the entire job is done. Sort out your thoughts; manufacture one key point for every section. If you make the four points listed above, you will need at least four paragraphs. Present surrounding information only when it is necessary to support a situation. The reviewer does not desire to read everything you know about a topic. Shape the theory specifically—do not take a broad view.

As always, give awareness to spelling, simplicity, and correctness of sentences and phrases.

Procedures (methods and materials):

This part is supposed to be the easiest to carve if you have good skills. A soundly written procedures segment allows a capable scientist to replicate your results. Present precise information about your supplies. The suppliers and clarity of reagents can be helpful bits of information. Present methods in sequential order, but linked methodologies can be grouped as a segment. Be concise when relating the protocols. Attempt to give the least amount of information that would permit another capable scientist to replicate your outcome, but be cautious that vital information is integrated. The use of subheadings is suggested and ought to be synchronized with the results section.

When a technique is used that has been well-described in another section, mention the specific item describing the way, but draw the basic principle while stating the situation. The purpose is to show all particular resources and broad procedures so that another person may use some or all of the methods in one more study or referee the scientific value of your work. It is not to be a step-by-step report of the whole thing you did, nor is a methods section a set of orders.

Materials:

*Materials may be reported in part of a section or else they may be recognized along with your measures.*

Methods:

- Report the method and not the particulars of each process that engaged the same methodology.
- Describe the method entirely.
- To be succinct, present methods under headings dedicated to specific dealings or groups of measures.
- Simplify—detail how procedures were completed, not how they were performed on a particular day.
- If well-known procedures were used, account for the procedure by name, possibly with a reference, and that’s all.

Approach:

It is embarrassing to use vigorous voice when documenting methods without using first person, which would focus the reviewer’s interest on the researcher rather than the job. As a result, when writing up the methods, most authors use third person passive voice.

Use standard style in this and every other part of the paper—avoid familiar lists, and use full sentences.

What to keep away from:

- Resources and methods are not a set of information.
- Skip all descriptive information and surroundings—save it for the argument.
- Leave out information that is immaterial to a third party.

Results:

The principle of a results segment is to present and demonstrate your conclusion. Create this part as entirely objective details of the outcome, and save all understanding for the discussion.

The page length of this segment is set by the sum and types of data to be reported. Use statistics and tables, if suitable, to present consequences most efficiently.

You must clearly differentiate material which would usually be incorporated in a study editorial from any unprocessed data or additional appendix matter that would not be available. In fact, such matters should not be submitted at all except if requested by the instructor.
Content:
- Sum up your conclusions in text and demonstrate them, if suitable, with figures and tables.
- In the manuscript, explain each of your consequences, and point the reader to remarks that are most appropriate.
- Present a background, such as by describing the question that was addressed by creation of an exacting study.
- Explain results of control experiments and give remarks that are not accessible in a prescribed figure or table, if appropriate.
- Examine your data, then prepare the analyzed (transformed) data in the form of a figure (graph), table, or manuscript.

What to stay away from:
- Do not discuss or infer your outcome, report surrounding information, or try to explain anything.
- Do not include raw data or intermediate calculations in a research manuscript.
- Do not present similar data more than once.
- A manuscript should complement any figures or tables, not duplicate information.
- Never confuse figures with tables—there is a difference.

Approach:
As always, use past tense when you submit your results, and put the whole thing in a reasonable order.

Put figures and tables, appropriately numbered, in order at the end of the report.

If you desire, you may place your figures and tables properly within the text of your results section.

Figures and tables:
If you put figures and tables at the end of some details, make certain that they are visibly distinguished from any attached appendix materials, such as raw facts. Whatever the position, each table must be titled, numbered one after the other, and include a heading. All figures and tables must be divided from the text.

Discussion:
The discussion is expected to be the trickiest segment to write. A lot of papers submitted to the journal are discarded based on problems with the discussion. There is no rule for how long an argument should be.

Position your understanding of the outcome visibly to lead the reviewer through your conclusions, and then finish the paper with a summing up of the implications of the study. The purpose here is to offer an understanding of your results and support all of your conclusions, using facts from your research and generally accepted information, if suitable. The implication of results should be fully described.

Infer your data in the conversation in suitable depth. This means that when you clarify an observable fact, you must explain mechanisms that may account for the observation. If your results vary from your prospect, make clear why that may have happened. If your results agree, then explain the theory that the proof supported. It is never suitable to just state that the data approved the prospect, and let it drop at that. Make a decision as to whether each premise is supported or discarded or if you cannot make a conclusion with assurance. Do not just dismiss a study or part of a study as "uncertain."

Research papers are not acknowledged if the work is imperfect. Draw what conclusions you can based upon the results that you have, and take care of the study as a finished work.
- You may propose future guidelines, such as how an experiment might be personalized to accomplish a new idea.
- Give details of all of your remarks as much as possible, focusing on mechanisms.
- Make a decision as to whether the tentative design sufficiently addressed the theory and whether or not it was correctly restricted. Try to present substitute explanations if they are sensible alternatives.
- One piece of research will not counter an overall question, so maintain the large picture in mind. Where do you go next? The best studies unlock new avenues of study. What questions remain?
- Recommendations for detailed papers will offer supplementary suggestions.
Approach:

When you refer to information, differentiate data generated by your own studies from other available information. Present work done by specific persons (including you) in past tense.

Describe generally acknowledged facts and main beliefs in present tense.

**The Administration Rules**

Administration Rules to Be Strictly Followed before Submitting Your Research Paper to Global Journals Inc.

*Please read the following rules and regulations carefully before submitting your research paper to Global Journals Inc. to avoid rejection.*

**Segment draft and final research paper:** You have to strictly follow the template of a research paper, failing which your paper may get rejected. You are expected to write each part of the paper wholly on your own. The peer reviewers need to identify your own perspective of the concepts in your own terms. Please do not extract straight from any other source, and do not rephrase someone else's analysis. Do not allow anyone else to proofread your manuscript.

**Written material:** You may discuss this with your guides and key sources. Do not copy anyone else's paper, even if this is only imitation, otherwise it will be rejected on the grounds of plagiarism, which is illegal. Various methods to avoid plagiarism are strictly applied by us to every paper, and, if found guilty, you may be blacklisted, which could affect your career adversely. To guard yourself and others from possible illegal use, please do not permit anyone to use or even read your paper and file.
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