



## Synthesis and Characterization of Nanocomposites Derived from Grafted Carboxymethyl Cellulose

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### Abstract

Nanoparticle (NP)-based polymer nanocomposites have received significant attention in recent years, particularly those based on carboxymethyl cellulose (CMC). This study aims to synthesize and characterize nanocomposites derived from grafted CMC. Because of the presence of polar functional groups, such as carboxyl and hydroxyl groups, in CMC, which contribute significantly to its strong adsorption capacity in metal structures, some research has indicated that this CMC is effective as a corrosion inhibitor, especially after being coated with NPs, as it protects metal surfaces from corrosion and helps them create a protective layer. In this study, nanocomposites derived from grafted CMC were synthesized. Initial CMC underwent a reaction with thionyl chloride in dimethylformamide (DMF) to produce CMC-Cl [I]. The CMC-Cl[I] reaction with various amines in DMF yields grafted CMC [II-X], which is subsequently blended with the natural biopolymer Chitosan to form blend polymers [XI-XIX]. Nanocomposites were synthesized through the interaction of blend polymers (grafted CMC/CS) with NPs such as magnesium oxide NPs (MgONPs), silver NPs (AgNPs), and gold NPs (AuNPs). The structures of the synthesized nanocomposites were characterized using FT-IR, <sup>1</sup>H-NMR, FESEM, and TEM. An investigation into the corrosion inhibition of blended polymers and nanocomposites on mild steel in 0.1 M HCl was conducted using weight-loss analysis. The results show that nanocomposites have a higher inhibition rate than blended polymers. Grafted CMC/CS/AuNPs [XXXI] reached an inhibition rate of 97% against the corrosion of carbon steel.

**Keywords:** Grafted carboxymethyl cellulose, Corrosion inhibition, Magnesium oxide nanoparticles, Nanocomposites.

### 1. Introduction

Carboxymethyl cellulose (CMC) is a cellulose derivative known to be the most common and widely used<sup>1</sup>. This is because its polar functional groups, such as carboxyl and hydroxyl groups, make it more effective at adhering to metal surfaces. People often use CMC in the food and pharmaceutical industries, but some research has shown that this cellulose derivative can also prevent corrosion<sup>2,3</sup>. It is a type of cellulose produced by combining sodium monochloroacetate with alkali cellulose. It is a polymer with several intriguing properties<sup>4</sup>; chemically stable, readily soluble, and non-toxic. It is also a biodegradable, inexpensive, biocompatible polymer that can form films<sup>5</sup>. The pharmaceutical, food, and packaging sectors frequently use it due to its safety and non-toxicity. CMC dissolves in water and acts as a stabilizer, making food-binding agents thicker<sup>6</sup>. Because CMC is a white, odorless, tasteless, and environmentally friendly water-soluble solid, it is used widely across many fields, including washing, paints, adhesives, textiles, paper, ceramics, food, and pharmaceuticals<sup>7</sup>. Because CMC has a higher viscosity, it adheres

better and acts as a stabilizer. This property makes it worthwhile in the food business as a thickening ingredient in ice cream, in medical goods as a coating for medicine capsules, and in pharmaceuticals as a gelling agent. On the other hand, it is excellent for the environment and can be modified chemically easily<sup>8-10</sup>. It has many hydroxyl and carboxyl groups in its structure, which makes it useful in many ways<sup>11</sup>. When CMC is combined with other polymers, such as CS. Chitosan is a natural polysaccharide and the second most common organic compound in nature, behind cellulose<sup>12</sup>. Chitosan is a weak base that is insoluble in water<sup>13</sup>. This feature is probably due to the strong hydrogen bonds that form between the chitosan molecular chains. It can, however, dissolve in weakly acidic water with a pH below<sup>14</sup>. When this happens, the glucosamine units ( $-NH_2$ ) transform into a form that can mix with water ( $NH_3^+$ ). The solubility of chitosan depends on where it comes from, how big its molecules are, and how much deacetylation it has<sup>15</sup>. Chitosan is a versatile chemical used in many fields, including medicine, agriculture, food stability, biocatalysis, and environmental applications. Chitosan has amino and hydroxyl groups that help it stick to metal surfaces, which is vital for its job as a corrosion inhibitor<sup>16</sup>. Over the past few years, biopolymers have been used to produce nanocomposites rapidly. Nanoparticles (NPs) such as gold, silver, and magnesium oxide are highly effective at inhibiting corrosion because they have a large surface area and are highly reactive<sup>17</sup>. Previous studies have demonstrated that NPs can prevent rusting<sup>18</sup>. Researchers are still looking at how to combine nanomaterials (NMs) with biodegradable polymers<sup>19</sup>. CMC is safe for living things, breaks down quickly, and is simple to find. CMC is often used to reduce and stabilize metal NPs (MNPs) because it is environmentally friendly and can be easily transformed into various forms. Nanoparticles improve the mechanical, thermal, biological, and barrier properties of biopolymers, which makes them more useful<sup>20</sup>. This study aims to synthesis and characterization of nanocomposites derived from grafted CMC.

## 2. Materials and Methods

### 2.1. Materials

Chemicals were provided by BDH, SCR, and CDH, gold NPs (AuNPs), silver NPs (AgNPs), and magnesium oxide NPs (MgONPs) by US, Research Nanomaterials, Inc

### 2.2. Instrumentation

The FT-IR spectra were recorded on a Shimadzu FTIR-8400S over the range  $400\text{ cm}^{-1}$  to  $4000\text{ cm}^{-1}$ . The  $^1\text{H}$ NMR spectra were obtained using an Ultra Shield 400 MHz spectrometer from Bruker, University of Tehran, Iran. TMS was used as an internal standard, with DMSO as the solvent. Field-emission scanning electron microscopy (FESEM) and transmission electron microscopy (TEM) were performed at the University of Tehran, Iran.

### 2.3. Methods of synthesis

#### 2.3.1. Synthesis of CMC-Cl [I]<sup>21</sup>

About 0.01 mol of CMC was combined with  $\text{SOCl}_2$  (0.01 mol) in 15 mL of MDF and refluxed for 8 hr. The excess thionyl chloride and MDF were removed under vacuum.

#### 2.3.2. Synthesis of grafted CMC [II-X]<sup>22</sup>

CMC-Cl [I] has been synthesised through the reaction of 0.1 g of CMC with 0.1 g of various amines, specifically 3,3'-dimethyl-[1,1'-biphenyl]-4,4'-diamine, 3,3'-dimethoxy-[1,1'-biphenyl]-4,4'-diamine, benzene-1,4-diamine, 4-nitroaniline, 4,6-dichlorobenzo[d]Thiazol-2-amine, sulfamethizole, trimethoprim, tacrine, 4-aminoantipyrine are combined in 20 mL of dimethylformamide (DMF), the mixture is refluxed for 8 hr. The solution was deposited onto a Petri dish and allowed to evaporate at ambient temperature.

#### 2.3.3. Synthesis of polymer blends [XI-XIX]<sup>23</sup>

Solvent casting was used to fabricate polymer mixes. The grafted CMC [II-X] was dissolved in water and stirred for 1 hr. at room temperature to prepare the grafted CMC solution. Additionally, 1 gram of chitosan was dissolved in 49 mL of a 2% aqueous acetic acid solution and stirred for 1 h. at room temperature to prepare the chitosan solution. Two polymer solutions,

grafted CMC [II-X] and chitosan, were blended in a 1:1 ratio to create a homogeneous solution using a hotplate stirrer for 1 hr following the amalgamation of the two polymer solutions.

#### 2.3.4. Synthesis of grafted CMC/CS nanocomposites [XX-XXXI]<sup>24</sup>

To help (magnesium, silver, or gold) NPs stick together in the composite matrix, 100 mg of dried grafted CMC/CS was added to 50 mL of MgONPs, AgNPs, or AuNPs solution at a concentration of 250 mg/L. For three hours, a hotplate stirrer was used to do this.

#### 2.3.5. Corrosion inhibition<sup>25</sup>

The chemical composition of carbon steel samples is as follows: C, 0.20; Fe, 99.23; Cu, 0.18; Si, 0.27; Mn, 0.05; Ca, 0.02; and S, 0.05. The gravimetric method, especially weight loss, is perhaps the most widely used method for measuring inhibition. The weight-loss method is easy to use and provides reliable results, making it the standard for measuring in various corrosion-monitoring systems. This study used 250 mL beakers containing 100 mL of the testing solution at room temperature, with total immersion, to measure weight loss. Weigh the iron coupons and hang them in a beaker by a rod and hook. The coupons were exposed for 6 hr, cleaned with distilled water, dried, and then weighed again. The weight loss, measured in grams, is the difference between the weight of the iron coupons before and after they were placed in different testing solutions. Using the following law, the inhibition efficiency % was calculated.

$$I.E. = (W_u - W_i / W_u) \times 100$$

$W_u$  stands for the weight loss in control solution,  $W_i$  represents the Loss in weight in the solution of inhibitor, I.E. stands for the Inhibition efficiency in inhibitor solution.

### 3. Results

The reaction of CMC with thionyl chloride in DMF made CMC-Cl [I]. The FTIR spectra of CMC-Cl showed a peak at  $1732\text{ cm}^{-1}$ , which is believed to come from the carbonyl group of acyl chloride. A grafted CMC was made by mixing CMC-Cl with various amines in DMF. The FT-IR spectra of polymer [II] have a strong peak at  $3367\text{ cm}^{-1}$ , showing that the OH stretching occurs because of hydrogen bonding. The little peaks at  $2924\text{ cm}^{-1}$  and  $2874\text{ cm}^{-1}$  are C-H stretch vibrations, and the band at  $1654\text{ cm}^{-1}$  comes from the amide group. The  $^1\text{H-NMR}$  spectrum of grafted CMC[II] shows a signal at  $\delta\ 8.13\text{ ppm}$  that comes from the amide proton ( $\text{C}=\text{O-NH}$ ). There are several peaks in the range of  $\delta\ (6.97\text{--}7.96)\text{ ppm}$ , which is where aromatic protons are found. A single signal at  $\delta\ (2.59\text{--}2.90)\text{ ppm}$  shows six protons from two  $\text{CH}_3$  groups. The peak that shows  $\text{CH}_2\text{COOH}$  on  $\text{C}_6$  of CMC is around  $(3.83\text{--}3.96)\text{ ppm}$ . The signals between 2.10 and 2.37 ppm are for ( $\text{H}_2, \text{H}_3, \text{H}_4, \text{H}_5$ ) of CMC, whereas the signals between 4.64 and 4.81 ppm are for  $\text{H}_1$  of CMC. Signals show the OH group of CMC at 5.96 ppm, and signals demonstrate the  $\text{CH}_2\text{COOH}$  group on  $\text{C}_6$  of CMC at  $12.6\text{ ppm}$ <sup>26,27</sup>.

Grafted CMC, the FT-IR results indicated that the blended polymer made from CMC and CS had a high level of miscibility. The broadening of the bands in the  $(2400\text{--}3600)\text{ cm}^{-1}$  range indicates that there is strong hydrogen bonding between the amino groups of CS and hydroxyl groups of CMC. The peak at  $1647\text{ cm}^{-1}$  for the ( $\text{C}=\text{ONH}$ ) functional group. Making grafted CMC/CS nanocomposites with MgONPs, AgNPs, and AuNPs. FT-IR data of nanocomposites [XX] show that the peaks at  $3288\text{ cm}^{-1}$  are due to O-H stretching caused by hydrogen bonds between and within molecules. The peaks at 2861 and  $2920\text{ cm}^{-1}$  are due to asymmetric and symmetric stretching vibrations of C-H from alkyl groups. The bonding of ( $\text{MgO}, \text{Ag}, \text{Au}$ ) NPs in the range of  $400\text{ to }800\text{ cm}^{-1}$  also supports the idea that particles are forming<sup>28,29</sup>.

#### 3.1. FESEM assay<sup>30</sup>

The surface morphology differs for grafted CMC, as illustrated in **Figure 1**. **Figure 2** depicts grafted CMC blended with CS, while **Figure 3** presents grafted CMC/CS/MgONPs. **Figure 4** showcases grafted CMC/CS/AgNPs, and **Figure 5** features grafted CMC/CS/AuNPs. Following the addition of CS to the grafted CMC, the results indicate alterations in the composite membrane's surface topography, significantly affecting cell spreading. The FESEM image

indicates a homogeneous dispersion of NPs over the matrix surface. The average particle size of grafted CMC [VIII] ranges from 642 to 782 nm. The average particle size of the grafted CMC blend with CS [XVII] ranges from 141 to 166 nm, while the average particle size of the grafted CMC/CS/MgO ranges from 48 to 80 nm in the presence of Mg. The average particle size of grafted CMC/CS/Ag ranges from 49 to 58 nm in the presence of silver, whereas the average NP size for gold NP ranges from 28 to 43 nm. The NPs exhibit a homogeneous distribution across the matrix surface. The particles in the nanocomposite film exhibited an almost spherical shape. Nonetheless, specific clusters of NPs were also observed, and the surface showed a degree of roughness. A uniform distribution of AuNPs in CMC/CS and robust NP adhesion to the polymer were demonstrated. The grafted CMC/CS mixture, together with NPs, forms new hydrogen bonds by disrupting the tight intramolecular hydrogen-bond network in CMC, thereby creating an adsorption site between the inhibitor and the metal.

### 3.2. TEM assay<sup>31-33</sup>

The TEM picture shows the improved CMC/CS solution, which was drop-cast with AgNPs and AuNPs. The (Ag, Au) NPs are spread out equally and are shaped like half-spheres. The particles are close together and point in opposite directions. Now that we have improved the CMC/CS, the AuNPs and AgNPs are more stable. The TEM image shows that the grafted CMC/CS/AgNPs and AuNPs are between 50 and 100 nm in size, as illustrated in **Figures 6** and **7**. Particles were spherical and had a thin layer of silver and gold around the grafted CMC/CS. TEM micrographs showed that the grafted CMC/CS was surrounded by layers of silver and gold nanostructures of uniform size. Gold and silver particles appeared to cover the grafted CMC/CS, leading to the growth of the CMC/CS/AgNPs and CMC/CS/AuNPs, as shown in **Tables 1-3** and **Schemes 1-3**.

**Table 1.** The FT-IR of polymers [II-X]

Com. No.	(O-H) of CMC	(C-H) aliph.	(C=O) amide	(C=C)	(C-O-C)
[II]	3367	2924,2874	1654	1590	1017
[III]	3345	2928,2867	1660	1579	1037
[IV]	3409	2930,2856	1680	1572	1033
[V]	3268	2922,2854	1647	1591	1023
[VI]	3306	3009,2869	1657	1594	1013
[VII]	4513	2930,2883	1661	1578	1030
[VIII]	3363	2935,2889	1651	1600	1035
[IX]	3317	2928,2835	1650	1591	1046
[X]	3316	2921,2870	1680	1584	1017

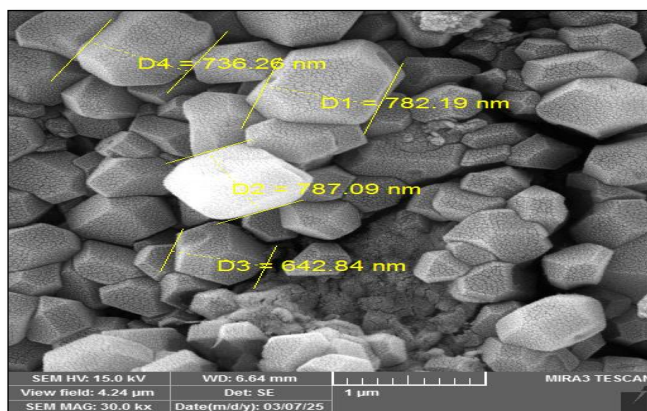
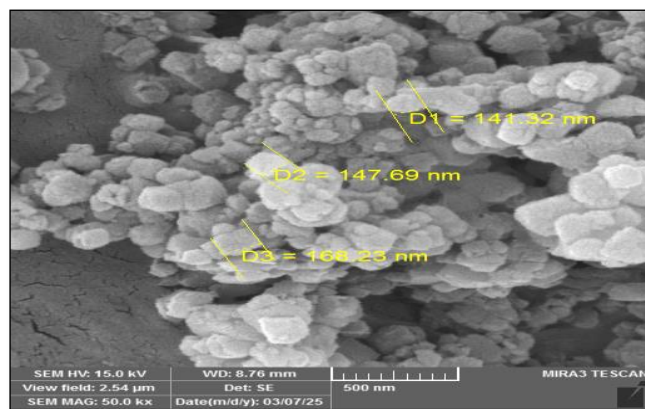
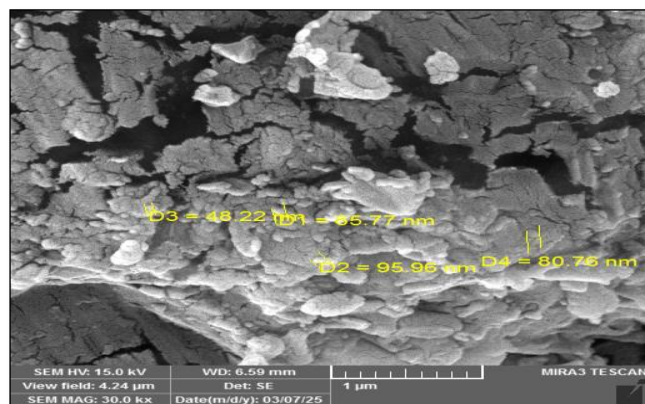
**Table 2.** The FT-IR of blend polymers [XI-XIX]

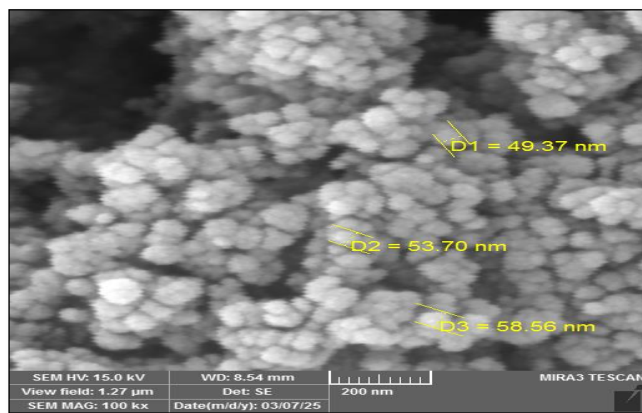
Com. No.	$\nu$ (O-H) and (N-H)	$\nu$ (C-H) aliph.	$\nu$ (C=O) amide.	$\nu$ (C=C) c	$\nu$ (-CH <sub>2</sub> -O-CO)	$\nu$ (C-O-C)
[XI]	3303	2973,2885	1647	1600	1274	1046
[XII]	3377	2921,2854	1648	1588	1241	1046
[XIII]	3356	2937,2885	1651	1601	1251	1051
[XIV]	3356	2925,2862	1640	1594	1255	1040
[XV]	3352	2915,2874	1648	1572	1255	1045
[XVI]	3341	2911,2874	1632	1564	1269	1021
[XVII]	3212	2920,2875	1640	1590	1259	1057
[XVIII]	3354	2918,2872	1640	1594	1251	1057
[XIX]	3350	2920,2874	1655	1585	1267	1058



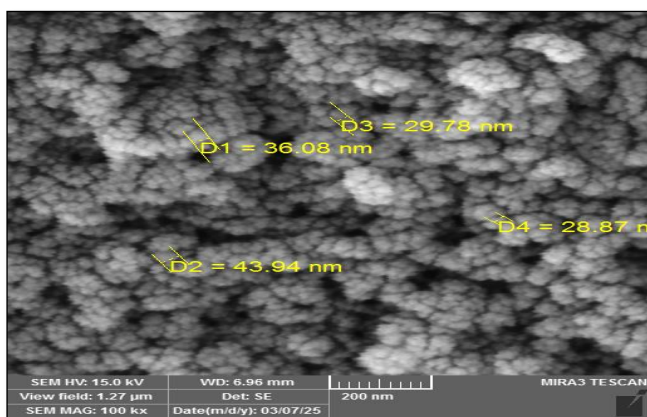
**Table 3.** Inhibition efficiency of blend polymer and nanocomposites in different concentrations of all polymers and nanocomposites at a temperature of room

Beaker No.	Compound	Conce. (ppm)	W <sub>1</sub>	W <sub>2</sub>	Loss in weight	%Loss in weight	I.E (%)
1	Control [HCl]	-	7.524	5.588	1.936	25.730	-
2	HCl+ grafted CMC/CS [ XVII]	100	7.293	7.012	0.281	3.853	85.485
3	HCl+ grafted CMC/CS [XVII]	200	7.754	7.495	0.259	3.340	86.621
4	HCl+ grafted CMC/CS /MgONPs [XXIX]	100	7.392	7.202	0.19	2.570	90.185
5	HCl+ grafted CMC/CS /MgONPs [XXIX]	200	7.761	7.590	0.171	2.203	91.167
6	HCl+ grafted CMC/CS /AgNPs[XXX]	100	7.495	7.384	0.111	1.480	94.266
7	HCl+ grafted CMC/CS /AgNPs[XXX]	200	7.593	7.489	0.104	1.369	94.628
8	HCl+ grafted CMC/CS /AuNPs[XXXI]	100	7.501	7.411	0.090	1.199	95.351
9	HCl+ grafted CMC/CS /AuNPs[XXXI]	200	7.453	7.414	0.039	0.523	97.985

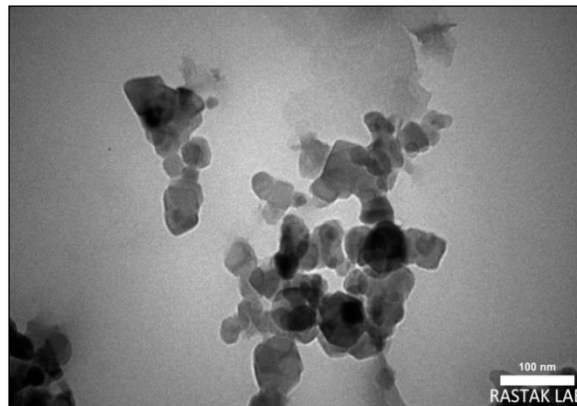
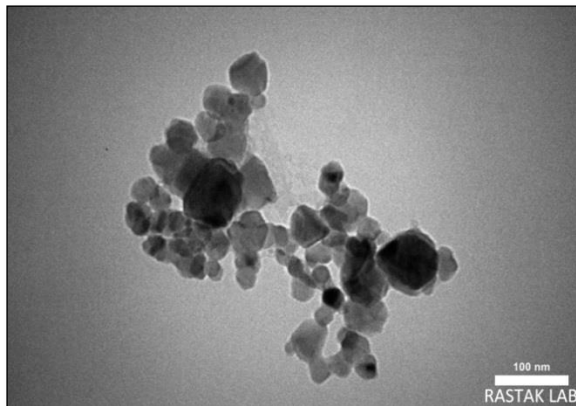
**Figure 1.** FESEM of grafted CMC[VIII]**Figure 2.** FESEM of grafted CMC/CS[XVII]**Figure 3.** FESEM of grafted CMC/CS/MgONPs[XXIX]



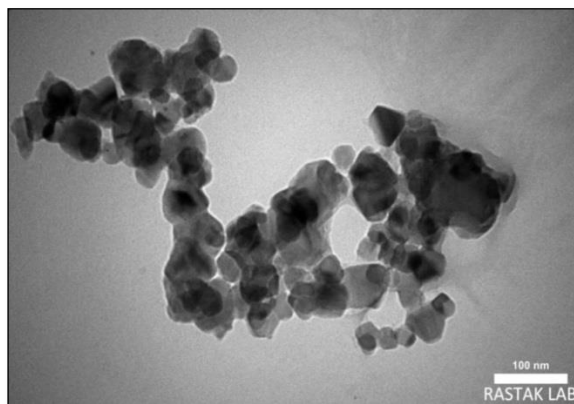
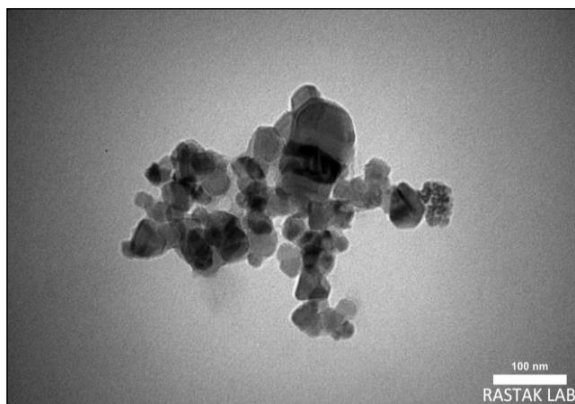
**Figure 4.** FESEM of grafted CMC/CS/AgNPs[XXX]



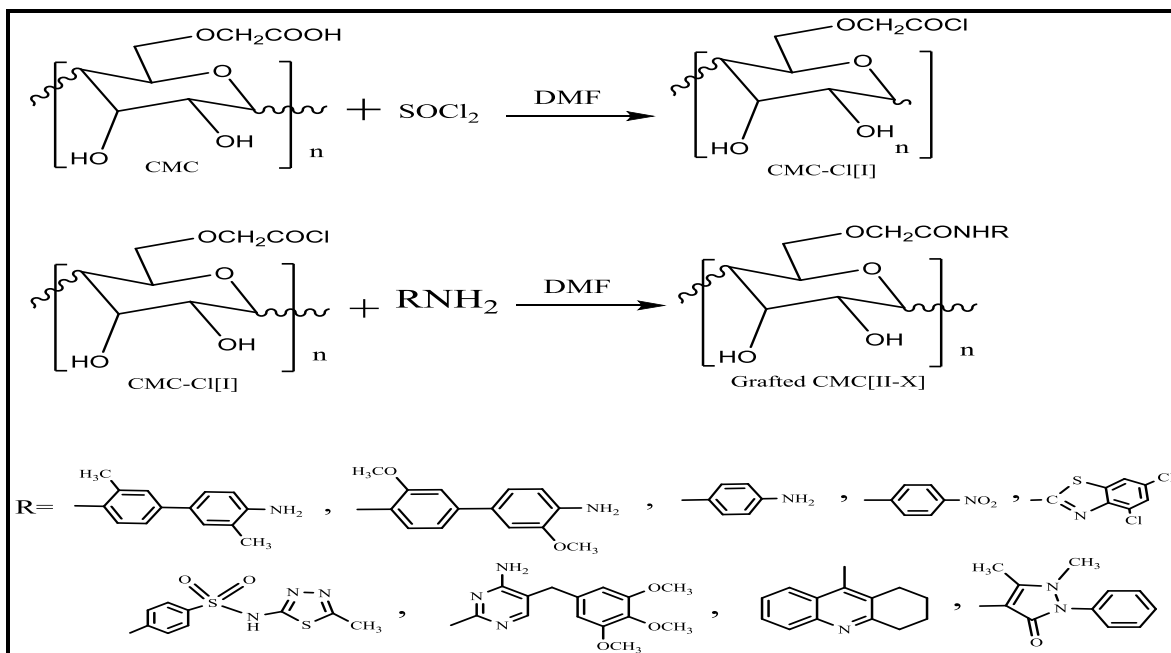
**Figure 5.** FESEM of grafted CMC/CS/AuNPs[XXXI]



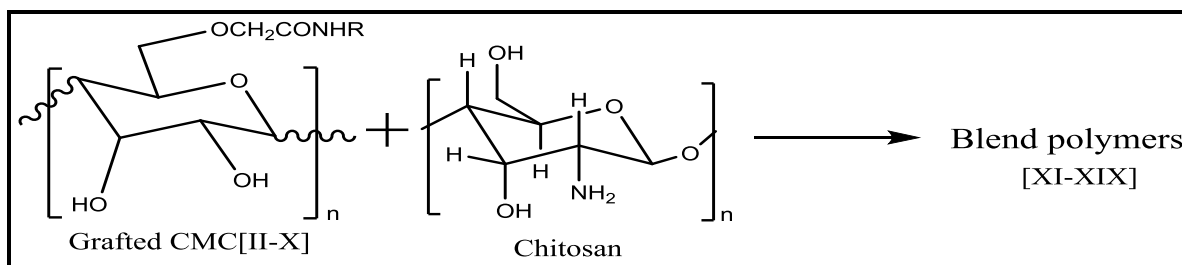
**Figure 6.** TEM of grafted CMC/CS/AgNPs



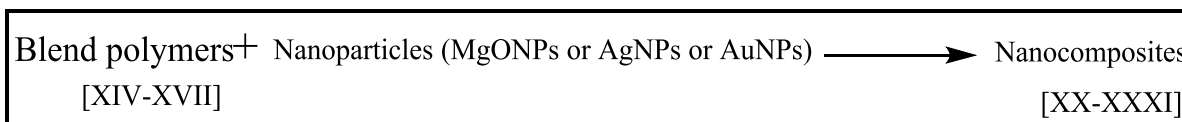
**Figure 7.** TEM of grafted CMC/CS/AuNPs



Scheme 1. Synthesis of grafted CMC [I-X].



Scheme 2. Synthesis of blend polymers [XI-XIX].



Scheme 3. Synthesis of nanocomposites [XX-XXXI].

#### 4. Discussion

Nanocomposites and their blend polymers have been shown to make carbon steel far less prone to corrosion in acidic environments. The azomethine group, electronegative atoms (N, S), and p electrons in aromatic systems are all reasons why this happens<sup>30</sup>.

The inhibitor compounds bind to the iron atoms via empty (d) orbitals, which are like spots that grab electrons. When the empty <sup>3</sup>d orbitals of Fe coincide with the p-orbital electrons in the inhibitor, they form coordinated bonds. The methoxy groups on the aromatic ring exert an inductive effect, making the ring more active and increasing electron density.

These modifications can make it much easier for the inhibitor to absorb, which boosts both protection and adsorption. The study indicates that compounds that adhere to the metal surface act as adsorption inhibitors, preventing corrosion. It's also important to note that NPs, CMCs, and CS compounds can enhance protection by forming a thin, inactive surface layer on the material. This approach keeps harmful chemicals from reaching the metal<sup>32,33</sup>.

This method either inhibits or slows redox processes in corrosion systems or renders dissolved oxygen ineffective. MgONPs, AgNPs, and AuNPs can protect metal surfaces against corrosion and form a protective layer. The results also showed that the nanocomposite grafted CMC/CS/AuNPs [XXXI] reached an inhibition rate of 97% against the corrosion of carbon steel<sup>34</sup>.

## 5. Conclusion

The current study seeks to synthesize nanocomposites via reactions initiated by altering CMC-Cl with different amines. Certain synthesized compounds and polymers exhibited satisfactory FT-IR, <sup>1</sup>H-NMR, FESEM, and TEM analyses that corresponded with data documented in the references. Finally, study the corrosion inhibition of blended polymers and nanocomposites in 0.1 M HCl on mild steel. The nanocomposites exhibit superior inhibitory efficiency compared to the blended polymer. The enhanced efficiency of the nanocomposites relative to the mixed polymer may be attributed to their NPs, resulting in the following efficiency hierarchy: [grafted CMC/CS/AuNPs [XXXI] > CMC/CS/AgNPs [XXX] > CMC/CS/MgONPs [XXIX] > CMC/CS [XVII]. The weight-loss data indicate that both surface coverage and inhibitory efficiency increase with increasing concentrations of the chemicals. The enhancement in inhibition efficiency with increasing inhibitor concentration may be attributed to the rise in –O–, –OH, and –COOH groups, which elevates the surface coverage of CMC molecules at higher concentrations.

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## Conflict of Interest

The authors have no conflict of interest.

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## Ethical Clearance

Procedures were conducted in accordance with ethical standards, and guidelines of the institution, the National Research Council, and national laws were followed.

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