Abstract

The class of imidazolium salts contains effective anticorrosion additives for metal substrates. This study evaluated the potential of 1-carboxymethyl-3-methylimidazolium bis (trifluoromethylsulfonyl) imide ($\text{HO}_2\text{CC}\text{1MImN}\text{TF}_2$) for application in cultural heritage, exploring it as anticorrosion additive in chitosan-based coatings for the protection of copper-based alloys. Under accelerated corrosion conditions with HCl vapor, the chitosan coating with $\text{HO}_2\text{CC}\text{1MImN}\text{TF}_2$ was less effective than the one with benzotriazole. The coating with a combination of $\text{HO}_2\text{CC}\text{1MImN}\text{TF}_2$ and benzotriazole resulted in the optimal protective efficacy of the bronze surface, and it also maintained high transparency without changing the bronze appearance.

Keywords: Biopolymer matrix, Anticorrosive, Copper-based alloy artifact, Ionic liquid

Introduction

Corrosion phenomena affecting metal surfaces exposed in polluted environments represent a serious global problem in several areas [1]. In the field of Cultural Heritage, for example, copper-based alloys have been widely used as raw materials by important artists in the last decades. Copper-based artworks, like other common metal objects, are subjected to corrosion processes due to the reaction of the metal surface with aggressive species present in the surrounding environment. The deterioration phenomena lead to the formation of both stable protective and reactive degrading species. Their chemical composition is mainly influenced by the composition of the alloy itself and by the environmental agents that come in contact with the object [1–5]. The corrosion products affect the aesthetical appearance of artworks and can compromise their structural properties [6].

Not only atmospheric pollutants, moisture or marine aerosol from outdoor environment can start the degradation processes. Aggressive species such as volatile organic compounds, particularly acetic and formic acid, or hydrogen sulphide, can be also present in indoor environments due to inadequate aeration, human emission, dust, construction materials and furniture.

For these reasons, metals substrates have to be protected to avoid degradation reactions and the formation of undesired products. Oxygen, humidity and chloride ions are particularly aggressive towards copper-based substrates since they induce a cyclic and irreversible degradation process known as “bronze disease”. It leads to the formation of corrosion products based on atacamite and its polymorphs ($\text{Cu}_2(\text{OH})_3\text{Cl}$), which are mainly responsible for the transformation of the alloy in a greenish powder. Moisture and oxygen cause also the formation of cuprite ($\text{Cu}_2\text{O}$) and the redness of the metallic surface. Sulphate-based pollutants promote the formation of...
basic sulphates such as brochantite CuSO₄·3Cu(OH)₃ or Cu₂SO₄·(OH)₆. Carbon dioxide and water, produce copper carbonate such as malachite Cu₂(OH)_2CO₃ and azurite Cu₃(OH)₂(CO₃)₂ [3].

Nowadays, organic coatings represent one of the most widespread approaches for the protection of metal substrates [7]. In the protection of bronze artworks, smart organic coatings have been widely proposed as a suitable approach for the inhibition of metal corrosion [8]. Such systems are characterized by a twofold protective action, as they are able to avoid the direct interaction of the metal surface with pollutants coming from the environment (passive protective mechanism) and to slow down degradation kinetics due to anticorrosive molecules that are released at the metal/coating interface and can interfere with the chemical degradation of metal (active protective mechanism). Nonetheless, protective coatings that are suitable for artistic objects should also meet some specific requirements such as transparency, long-term efficiency and reversibility or, at least, re-applicability. Generally, the commonly used commercial products are based on acrylic resins or microcrystalline waxes dispersed in harmful solvents such as toluene and white spirit. Some notable examples include Incralac®, which consists of Paraloid B44™, a copolymer of methyl methacrylate with an undefined monomer, dispersed in toluene, and Soter, microcrystalline waxes dispersed in a turpentine and ether mixture. Often these products are combined in double-layer systems to enhance their protective actions (such as Incralac® and wax) [9]. Fluoropolymers have also been used because of their protective actions (such as Incralac® and wax) [9].

In this perspective, BTA-functionalized imidazolium liquids have been synthesized as effective corrosion inhibitors for steel and copper [29]. Their corrosion inhibition mechanism is similar to what is known for most of the organic corrosion inhibitors, blocking the anodic and cathodic reactive sites on a metallic surface through: (a) electrostatic adsorption of IS ions on anodic and cathodic surface sites; (b) chemisorption; and (c) surface interactions between IS compounds, forming a 2D barrier coating [30, 31].

The present work fits in this frame; focused on the use of IS to improve the performances of CHT-based coatings intended for the protection of bronze surfaces. More specifically, we investigated the potential of the carboxyl-functionalized IS 1-carboxymethyl-3-methylimidazolium bis (trifuoromethysulfonyl) imide (HO₂CC₂MImNTf₂; Fig. 1) as additive in CHT-based coatings for the inhibition mechanism is similar to what is known for most of the organic corrosion inhibitors, blocking the anodic and cathodic reactive sites on a metallic surface through: (a) electrostatic adsorption of IS ions on anodic and cathodic surface sites; (b) chemisorption; and (c) surface interactions between IS compounds, forming a 2D barrier coating [30, 31].

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In this context, imidazolium salts (IS), including ionic liquids that are IS with a melting point of 100 °C or lower, possess favorable characteristics for their application in formulations for protective coatings, e.g. negligible vapor pressure, chemical and thermal stability and low flammability [26, 27]. Generally, they consist of an imidazolium cation and an organic or inorganic anion, and their properties can be tuned by structural changes in the ions, enabling the preparation of task-specific ionic liquids [28]. Besides, their negligible vapor pressures turns them into ideal candidates to minimize the risk of inhalation. In this perspective, BTA-functionalized imidazolium ionic liquids have been synthesized as effective corrosion inhibitors for steel and copper [29]. Their corrosion inhibition mechanism is similar to what is known for most of the organic corrosion inhibitors, blocking the anodic and cathodic reactive sites on a metallic surface through: (a) electrostatic adsorption of IS ions on anodic and cathodic surface sites; (b) chemisorption; and (c) surface interactions between IS compounds, forming a 2D barrier coating [30, 31].

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![Fig. 1 Imidazolium salt explored in this study](image-url)
exploitation of several roles: (a) anticorrosion additive; and (b) co-anticorrosion additive in the presence of BTA. In particular, \( \text{HO}_2\text{CC}_1\text{MImNITf}_2 \) was selected for the development of coating formulations, showing benefits over the IS presented in Fig. 2. The application of coatings with hydrophobic IS (Fig. 2a) protected the alloy only in the areas covered with IS. Most likely the incompatibility between the hydrophobic IS and the hydrophilic CHT caused the presence of uncovered areas. Similarly, BTA-functionalized IS (Fig. 2b) did not afford the desired protection due to their low solubility’s in water-ethanol (50:50), which was used to solubilize CHT. Within the screened hydrophilic IS (Figs. 1 and 2c), which are soluble in water-ethanol (50:50), \( \text{HO}_2\text{CC}_1\text{MImNITf}_2 \) was the most promising IS. The optimum \( \text{HO}_2\text{CC}_1\text{MImNITf}_2 \) content was chosen to be 35 wt% with respect to the CHT content. Besides, 2 wt% of BTA, with respect to the CHT content, was selected, being comparable with the amount typically used in commercial products.

The protective efficacy of the investigated IS formulations was evaluated by submitting coated bronze disks to an accelerated test in acidic conditions. In order to promote the corrosion processes at the alloy surface, the disks were treated with acidic water vapors containing chloride ions by using a procedure previously reported [14, 15, 32–34]. The degradation processes in bronze objects are very complex and can be induced by several species. Therefore, chloride ions were selected to perform the tests since they are among the most aggressive degrading agents for copper-based alloys. In this way, it was possible to rapidly obtain information about the protective efficacy of the investigated formulations.

A copper-based alloy, with chemical composition and dendritic metallurgical structure similar to those commonly used in bronze objects of art, was used as the metallic substrate. In this work the comprehensive results related to the morphological, chemical and functional characterization of the best CHT-based coatings modified with BTA, \( \text{HO}_2\text{CC}_1\text{MImNITf}_2 \) and the mixture BTA- \( \text{HO}_2\text{CC}_1\text{MImNITf}_2 \) are reported. The results confirm that the selected IS, \( \text{HO}_2\text{CC}_1\text{MImNITf}_2 \), does not affect the esthetical characteristics of pristine CHT-based coating and, when used at 35 wt% with respect to CHT and in the presence of 2 wt% of BTA, has a positive effect on the anticorrosion effectiveness of the resulting coating when compared to the CHT system containing only 2 wt% of BTA. This result is likely ascribed to an improvement of interfacial adhesion between the CHT matrix and the metal substrate. To the best of our knowledge, this preliminary study on the active protection of bronze alloys with CHT-based coatings modified with BTA and the IS \( \text{HO}_2\text{CC}_1\text{MImNITf}_2 \) is innovative and similar uses of IS in cultural heritage were not reported so far.

**Experimental**

**Materials**

1-Methylimidazole (99%), methyl 2-chloroacetate (99%), lithium bis (trifluoromethylsulfonyl) imide (LiNTf2, 99.95%), dichloromethane (P.A.), magnesium sulfate (≥ 99.5%), silver nitrate (99.5%), CHT (medium molecular weight, viscosity 200–800 cP, 75–85% deacetylated), D-(+)-gluconic δ-lactone (DGL, ≥ 99.9%), ethanol (≥ 99.8%), water Cromasolv plus for HPLC, hydrochloric acid (37%, ACS reagent) and sodium hydroxide (≥ 97.0%, pellets, ACS reagent) were purchased from Sigma-Aldrich. BTA (99%) was purchased from Bresciani. Glacial acetic acid (99.9%) was purchased from Carlo Erba. Before using, CHT was washed in boiling water for 1 h, filtered, thoroughly washed with distilled water to remove impurities, and dried under vacuum for 12 h.

**Bronze substrate**

A copper-based alloy with nominal composition of 85 wt% Cu, 5 wt% Sn, 5 wt% Pb, and 5 wt% Zn, produced by casting and with a dendritic structure, was used as representative metal substrate [3]. The alloy disks have been polished by using SiC papers at 1200 grit and, subsequently, diamond pastes up to 1/4 μm. After polishing, the Cu-based alloy disks were cleaned with ethanol.
Atomic force microscopy (AFM)
Surface roughness was measured with a Dimension 3100 atomic force microscope, equipped with a NanoScopeIV controller (Veeco, Santa Barbara, CA), operating in tapping mode. Phosphorus-doped silicon probes with a resonant frequency of around 275 kHz and a nominal spring constant of 20/80 Nm were employed (RTESP) (Veeco, Santa Barbara, CA). A scan rate of 0.2 Hz was employed at a resolution of 384 pixels/line. The root mean square roughness of polished bronze substrate was determined in a scan area of $50 \times 50 \mu m^2$.

Synthesis of 1-carboxymethyl-3-methylimidazolium bis(trifluoromethylsulfonyl) imide ($HO_2CC_1MImNTf_2$)
Scheme 1 shows the synthetic route that was applied for the preparation of $HO_2CC_1MImNTf_2$. Initially, a literature procedure was used for the synthesis of 1-carboxymethyl-3-methylimidazolium chloride ($HO_2CC_1MImCl$) [35]. A bimolecular nucleophilic substitution reaction (SN$_2$) between 1-methylimidazole (2.02 g, 24.6 mmol, 1.00 equiv.) and methyl 2-chloroacetate (2.30 g, 24.6 mmol, 1.00 equiv.) was performed for 1 h at 0°C to prepare the ester-functionalized IS 1-methoxycarbonylmethyl-3-methylimidazolium chloride ($H_3CO_2C_1MImCl$). Subsequently, this IS was treated with HCl (24.6 mmol, 1.00 equiv.) for 3 h at 100°C to synthesize $HO_2CC_1MImCl$.

A literature procedure was used for the anion metathesis [36]. $HO_2CC_1MImCl$ (0.503 g, 2.85 mmol, 1.00 equiv.), LiNTf$_2$ (0.858 g, 2.99 mmol, 1.05 equiv.) and water (25.0 mL) were mixed and stirred at room temperature (rt) for 24 h. This solution was extracted with dichloromethane (3 x 35 mL), and the joined organic phase was dried over magnesium sulfate. Filtration followed by solvent evaporation afforded $HO_2CC_1MImNTf_2$.

Nuclear magnetic resonance (NMR)
NMR spectra of $HO_2CC_1MImCl$ and $HO_2CC_1MImNTf_2$ were recorded on Bruker (400 MHz) and Varian Inova (300 MHz) spectrometers at ambient temperature. The chemical shifts are given in parts per million (ppm) and referenced to the residual solvent signal ($D_2O = 4.79$ ($^1H$); dimethyl sulfoxide-$d_6 = 39.52$ ($^{13}C$)).

In vitro cytotoxicity of $HO_2CC_1MImNTf_2$ and BTA
For the cytotoxicity tests, IS and BTA were washed several times with phosphate buffer solution and freeze dried to obtain dry powders. Then, IS and BTA solutions in Dulbecco’s modified Eagle’s medium (GIBCO® by Life Technologies, Italy) at concentrations of 10, 100, 200, 350 and 500 $\mu g \cdot mL^{-1}$ were prepared in an incubator shaker at 37°C for 3 h, and then sterilized for 2 h under ultraviolet

![Scheme 1 Synthetic route for $HO_2CC_1MImNTf_2$](image-url)
radiation ($280 < \lambda < 200$). The wells of Falcon® 48-well culture plates were seeded with 5000 cells (50 μL) of 80% confluent mouse fibroblast (L929) cell line (in accordance with the standard international guideline ISO10993-5), followed by the addition of 350 μL of a chosen IS or BTA solution. Dulbecco's modified Eagle's medium without IS and BTA was used as negative (non-toxic) control. The incubation was performed at 37 °C (5% CO₂, 97% air humidity) for 1 and 5 days of exposure. Alamar Blue (Invitrogen, Italy) was used as a redox indicator to evaluate biocompatibility. This assay quantified the redox indicator that changed to a fluorescent product in response to chemical reduction by mitochondrial enzymes such as dinucleotide adenine flavin dehydrogenase and dinucleotide adenine nicotinamide dehydrogenase, providing a quantifiable indication for the metabolic activity of living cells. After the selected times of 1 and 5 days, the medium was removed and a 200 μL aliquot of diluted Alamar Blue (1:10) in red phenol-free medium (GIBCO® by Life Technologies, Italy) was added to each well. After incubation for 4 h at 37 °C (5% CO₂, 97% air humidity), 100 μL of each solution was transferred to a 96-well plate for colorimetric analysis ($n = 4$). Wells without any cells were used to correct any background interference of the redox indicator. Cell viability was correlated with the magnitude of the dye reduction and expressed as % AB reduction according to the manufacturer’s protocol.

Preparation of chitosan-based coatings and characterization

The pure CHT formulation (CHT) was prepared by adding the CHT powder (1.0 wt%/vol%) to an aqueous DGL solution (0.05 M) and then the mixture was kept under stirring for 24–30 h to achieve complete solubilization of the polymer. The final pH of the solution was adjusted to 6 by the addition of 1 M sodium hydroxide solution. The corrosion inhibitor BTA (2.0 wt% with respect to CHT) was added to the CHT suspension from DGL (pH = 6) to prepare the formulation CHT/BTA. The formulation CHT/IS was prepared by dissolving 35 wt% (with respect to CHT) of HO₂CC₂MImNTf₂ in the water/ethanol CHT solution from DGL (pH = 6) to prepare the formulation CHT/IS. The formulation CHT/BTA-IS was prepared by adding both BTA (2.0 wt% with respect to CHT) and HO₂CC₂MImNTf₂ (35 wt% with respect to CHT) to the pure CHT solution from DGL (pH = 6). All the solutions were diluted with ethanol to obtain formulations with a final water/ethanol composition of 50/50 vol%/vol%. The coatings were prepared by solvent casting, dropping 320 μL of the water/ethanol CHT-based formulations on the reference bronze disks with a diameter of 30 mm and subsequent drying overnight at rt.

Colorimetric analysis

Color differences between an uncoated copper-based alloy disk and disks coated with CHT, CHT/BTA, CHT/IS, or CHT/BTA-IS were measured with an Exact color spectrophotometer (X-Rite, Grand Rapids, MI, USA). Color differences in the CIELab space were evaluated by using the uncoated copper disk as a reference and measuring the $\Delta L^*$, $\Delta a^*$ and $\Delta b^*$ values, where $L^*$ is the color lightness, $a^*$ is the green-magenta contribution and $b^*$ is the blue-yellow contribution. For each sample, three colorimetric measurements were performed and average values of $\Delta L^*$, $\Delta a^*$, and $\Delta b^*$ were obtained. For each couple of specimens, the Euclidean difference $\Delta E^*$ was calculated using Eq. 1 [37]:

$$\Delta E = \sqrt{(\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2}$$

(1)

Nanoscratch test

The nanoscratch test was performed to investigate the abrasion resistance of the coatings. The experiments were carried out with a NanoTest Platform made by Micro Materials Ltd., equipped with a Berkovich diamond tip of three-side pyramidal geometry. In a typical experiment, the indenter tip was moved tangentially over the coating surface at a fixed velocity of 1 μm s⁻¹ while progressively increasing the applied normal load up to 50 mN over a scratch length of 2 mm.

Accelerated corrosion test

In order to assess the protective efficacy of the CHT-based coatings, accelerated corrosion tests were carried out by heating the coated bronze disks inside a closed glass vessel at 50 °C in the presence of HCl vapor (aqueous 1.0 M HCl solution) and 100% relative humidity (RH). Note that bronze disks were exposed to acid vapor but not in direct contact with the HCl solution (Fig. 3).
The coated bronze disks were characterized before and after the test at different time intervals (up to 12 h). All the characterizations were conducted in the central part of the disks where the border effect is not relevant. After the corrosion treatment, the CHT-based films were partially removed by using tissue paper soaked in water and ethanol.

Optical microscopy (OM) and image analysis
For the evaluation of the protective efficacy of the coatings, modifications at the metal surface were analyzed by means of OM. The investigations were carried out with a Leica MEF 4 M microscope equipped with a digital camera (Leica DFC 280). Image analysis has been exploited to characterize the state of the metal surface after the accelerated corrosion test. OM images at the metal/coating interface were collected without removing the coating, and the percentage of corroded surface was estimated through a custom-tailored image analysis protocol, whose details can be found elsewhere [14, 15]. Briefly, OM images were first equalized and converted into grayscale, in which lighter tones correspond to not corroded areas, while darker ones to corroded areas. Then, a threshold tone distinguishing corroded and not corroded areas was arbitrary and uniquely defined. Finally, the percentage of corroded area was estimated from the histograms corresponding to the relative frequency of the tones extracted from each grayscale image. Images of untreated coated disk were used as reference.

Attenuated total reflectance–Fourier transform infrared (ATR–FTIR) spectroscopy
Bronze disks with CHT-based coatings containing BTA and/or HO2CC1MImNTf2 were characterized before and after the accelerated corrosion test by ATR–FTIR spectroscopy. This was performed with the aid of a Nicolet iS50 spectrometer (Thermo Fisher Scientific). The samples were characterized by forcing their coated surface in contact with the diamond crystal and taking care that the pressure was constant for all analyses. The spectra were collected by 32 scans at a resolution of 4 cm⁻¹. All the spectra were then normalized with respect to the peak a 1583 cm⁻¹, which can be ascribed to the N–H bending of the CHT primary amine groups and the DGL used for the CHT solubilisation in water [14, 38]. No ATR correction has been applied to the data.

Scanning electron microscopy (SEM) analysis
The determination of the thickness (before and after the corrosion test), and micro-chemical and -morphological characterisations of the coatings (before and after the corrosion test) were performed by means of a scanning electron microscope Cambridge 360 equipped with a LaB6 filament and a high brilliance LEO 1530 field emission scanning electron microscope apparatus equipped with an energy dispersive X-ray spectrometer (EDS).

Results and discussion
HO2CC1MImCl (Scheme 1) was obtained as white solid and the two-step synthesis resulted in 92% yield. The recorded ¹H (Additional file 1: Figure S1) and ¹³C NMR (Additional file 1: Figure S2) spectra are in agreement with the data published previously [35]. As chloride anions promote the bronze disease, anion methatesis between chloride and bis(trifluoromethylsulfonyl) imide was performed (Scheme 1). This reaction resulted in the formation of HO2CC1MImNTf2 as transparent oil in 38% yield. A test with acidified silver nitrate indicated the absence of chloride. The ¹H (Additional file 1: Figure S3) and ¹³C NMR (Additional file 1: Figure S4) spectra correspond to the chemical structure of HO2CC1MImNTf2. When considering the ¹H NMR data (300 MHz, D₂O): δ (ppm) 8.74 (s, 1H, C2-H), 7.45–7.44 (m, 2H, C4-H and C5-H), 5.02 (s, 2H, CH2), 3.89 (s, 3H, CH3); the number of peaks and their corresponding integrals verify the chemical identity of HO2CC1MImNTf2. The same counts for the ¹³C NMR data (100 MHz, dimethyl sulfoxide-d6): δ (ppm) 169.8 (CO2H), 136.7 (C2), 122.9 (C4/5), 122.8 (C5/4), 118.6 (qua, J = 319.7 Hz, CF₃), 49.6 (CH₃), 35.2 (CH₃). MS-ESI(+) detected m/z 141.1201 for the cation of HO2CC1MImNTf2, which is close to the calculated value of 141.0659 for C₆H₉N₂O₂⁺.

For HO2CC1MImNTf2 and BTA, their cytotoxicity were determined in cells from the mouse fibroblast cell line (Fig. 4). Compared to the cell proliferation of the control (IS- and BTA-free), HO2CC1MImNTf2 showed similar cell viability after 1 day, independent of the applied IS concentration. After 5 days, the cell proliferation in the presence of HO2CC1MImNTf2 was without significant difference compared to the control or about 30% lower for the samples at 200 and 500 μg mL⁻¹ (Fig. 4a). Overall, the obtained cytotoxicity data on in vitro murine fibroblast cells indicated no cytotoxic effect of HO2CC1MImNTf2 and the safe use of this IS in the protective formulations as demonstrated by the capability of HO2CC1MImNTf2 to support increasing cell proliferations over the culture time. BTA showed a different profile compared to the control or about 30% lower for the samples at 200 and 500 μg mL⁻¹ (Fig. 4b). Now, the cytotoxicity increased with increasing BTA concentrations. At the lower concentrations of 10 and 100 μg/mL, although BTA was not toxic after 1 day, a significant reduction in cell proliferation (about 50% compared to the control) was observed at day 5, where the toxic effect was evident. These results confirm that HO2CC1MImNTf2 is less cytotoxic than BTA, which is an important insight
that justifies the strategic selection made in this paper to address the research towards more sustainable anticorrosive compounds as compared with traditional BTA.

To assess the efficacy of the CHT-based coatings functionalized with the IS \( \text{HO}_2\text{CC}_1\text{MImNTf}_2 \) and/or BTA, a copper-based alloy was used as metal model substrate. The alloy disks were polished to obtain a flat and smooth surface with a mirror-like finish. Surface analysis by AFM enabled the determination of the root mean square roughness, being \( 17.7 \pm 4.31 \) nm. The preparation of CHT-based formulations was optimized to obtain uniform, compact and transparent films after deposition on Cu-based disks and drying. In particular, DGL was used to solubilize the polymer and an ethanol/water mixture with composition of 50/50 vol%/vol% was selected as liquid medium (formulation CHT). The commonly used corrosion inhibitor BTA (2 wt% with respect to CHT; formulation CHT/BTA), the IS \( \text{HO}_2\text{CC}_1\text{MImNTf}_2 \) (35 wt% with respect to CHT; formulation CHT/IS) and the mixture of BTA and \( \text{HO}_2\text{CC}_1\text{MImNTf}_2 \) (2 and 35 wt% with respect to CHT, respectively; formulation CHT/BTA-IS) were added to the CHT formulation and investigated as protective agents.

The colour coordinates for the different samples and the \( \Delta E^* \) values calculated with respect to the pristine bronze disk are reported in Table 1. Colorimetric analysis in the CIELab space revealed that the coatings present a very small impact on the aesthetic appearance of the original bronze substrate. The \( \Delta E^* \) values, indeed, are small and compatible for applications in cultural heritage, which is \( \Delta E^* \) below the threshold of 5, being considered the lower limit for human eye perception [39, 40]. The only exception is represented by the coating obtained from the formulation CHT/BTA, which presented a significant variation with respect to the pristine bronze disk (i.e. \( \Delta E^* > 5 \)). Looking at the color coordinates, it emerges that the largest variation with respect to the uncoated bronze surface was found for the \( L^* \) value (variation of \(-38\%)\). This indicated that the presence of the CHT/BTA coating caused a significant reduction of the color lightness of the metal substrate, probably due to a poor dispersion of BTA in the chitosan matrix and/or the formation of small BTA crystalline precipitates. At the same time, we highlight that the coating containing both the IS and the corrosion inhibitor (CHT/BTA-IS) exhibited the lowest \( \Delta E^* \) among the coatings with additives, its value being very similar to that related to the CHT coating. This suggested a synergic effect between IS and BTA in not affecting the appearance of the metal surface. In this sense, it is worth noting that the \( \Delta E^* \) values of the coatings developed in the present work, and in particular that of the CHT/BTA-IS coating, are much lower than other values reported in the literature for BTA-containing protective coatings for bronze surfaces [41].

Figure 5 shows the ATR–FTIR transmittance spectra of the prepared coatings on bronze disks. The spectrum of the CHT coating exhibits the typical peaks of DGL.
and CHT polymer according to the assignments already reported [14]. In the case of CHT/BTA, the BTA content of 2 wt% with respect to the amount of polymer was too low to be detected in the spectrum. On the other side, the presence of IS \( \text{HO}_2\text{CC}_1\text{MImNTf}_2 \) at 35 wt% with respect to the amount of polymer was clearly evidenced in the spectrum of CHT/IS by three transmittance peaks in the range of 1400–1100 cm\(^{-1}\). It is worth noting that one of these peaks is clearly observed in the spectrum of pristine \( \text{HO}_2\text{CC}_1\text{MImNTf}_2 \) at 1358 cm\(^{-1}\) and may be ascribed to C–N (stretching vibration) bands in imidazolium rings [42]. The other two peaks at around 1220–1230 cm\(^{-1}\) may tentatively be ascribed to intimate CHT-IS interactions. Part of the CHT amines is protonated (being solubilized in acidic conditions), which may interact with the IS anion, bis (trifluoromethylsulfon) imide.

The coating thickness measured by SEM was about 8–14 \( \mu \text{m} \) for all the investigated samples (Additional file 1: Figure S5). SEM micrographs showed that the surfaces were homogeneous without apparent cracks or domains due to separations of components. A representative SEM micrograph is reported in Fig. 6, showing in the right part the surface coated with CHT/BTA-IS and in the left part the bare surface of bronze disk after removal of this coating. The EDS spectra confirmed the effective removal of the coating (Fig. 6). At higher magnification, the surface of the coating appears without any cracks and any phase separation confirming that all components of the coating are fully miscible in the composition range investigated.

A representative output of the progressive-load nanoscratch measurements is given in Fig. 7, showing both the optical micrographs of the scratch paths and the initial parts of the nanoscratch profile curves, which enabled determining the coating failure.

Nanoscratch tests offer the possibility of assessing the quality of the coating/substrate interactions. This can be done by looking at the coating failure mode during/after scratch and also by comparing the depth vs. load curves.
recorded during the scratch [43]. The analysis of the optical micrographs of the scratch paths in Fig. 7 highlights that a modest deformation occurred for relatively low loads for all the coatings. On the contrary, at higher loads the formation of pile-up material on either side of the track becomes significant, the effect being much more pronounced in the case of samples containing BTA. The latter presumably plasticizes the polymer matrix [14], which on the contrary showed a very good adhesion to the metal surface. More specifically, when the coating/substrate adhesion is poor, a significant amount of coating material is generally stripped off along the entire length of the scratch path [44]. This is the case of the CHT/BTA sample. On the contrary, the addition of the IS only slightly affected the already good chitosan-bronze adhesion, as indicated by the similarity of the scratch paths of pristine polymer (CHT) and polymer with IS (CHT/IS). From a practical perspective, a comparative evaluation of the coatings can be done in terms of critical scratch load, which is the load value at which the coating fails [45]. In this sense, the CHT coating exhibited the best performances, followed by CHT/IS, CHT/BTA-IS, and CHT/BTA. This rank is further supported by the profiles of the scratch curves, especially when looking at their initial part (i.e. up to 30 mN of applied load). Besides the different slope, which is mainly related to the different grade of flatness of the bronze substrates, the curves in Fig. 7 are clearly rougher for the samples containing BTA (both CHT/BTA and CHT/BTA-IS). This indicates that its addition caused a lowering of the abrasion resistance of the CHT matrix and decreased the adhesion of the coating to the underlying metal surface. Nonetheless, a closer look at the curves also suggests that the presence of the IS in CHT/BTA-IS helped in partially recovering the good adhesion to the substrate and high scratch resistance of the pristine CHT coating. This behaviour can be possibly ascribed to the electrostatic interactions that took place between the metal cations present on the bronze surface and the IS bis (trifluoromethylsulfonyl) imide anions (Fig. 1), abbreviated as NTf₂:

\[
M^{n+} + n(\text{NTf}_2^-) \rightarrow \text{M}^{n+} \cdots n(\text{NTf}_2^-)_{\text{ads}}
\]

Due to this interaction, the positively charged CHT matrix (rich of ammonium ions due to the solubilisation performed in acidic conditions) may strongly compete with the imidazolium cations of the IS to interact with the functionalized substrate. As a consequence, the polymer forms a continuous film that protects the substrate and increases the adhesion to the metal of the modified film:

\[
\text{M}^{n+} \cdots n(\text{NTf}_2^-)_{\text{ads}} + \text{Chit}^{n+} \rightarrow \text{M}^{n+} \cdots n(\text{NTf}_2^-)_{\text{ads}} \cdots \text{Chit}^{n+}
\]

Although further investigations are necessary, a similar mechanism was recently considered to explain the protective effect of dicationic IS on steel substrates in acidic conditions [46].

With the aim to investigate the protective efficacy of the developed coatings, all the coated disks were subjected to accelerated corrosion treatments and the occurrence of surface modifications was observed at different time intervals by optical microscopy.

In the case of the coatings CHT and CHT/IS, significant modifications of the alloy surface were observed after 6 h of accelerated corrosion test (Fig. 8).

These results suggest that the CHT and CHT/IS coatings were not stable and able to prevent the occurrence of corrosion processes under the tested conditions. The hypothesis of the coating degradation was supported by ATR–FTIR analysis since the small peak at 880 cm⁻¹, typical of the wagging (the CH bending out of plane of the saccharide ring) of the CHT structure
disappeared after the exposure to aggressive vapors. Moreover, the ATR–FTIR spectra of the CHT and CHT/IS coatings, after exposure to HCl vapors, revealed the presence of two weak bands in the region between 1000 and 800 cm\(^{-1}\), being characteristic for the bending of the hydroxyl groups in Cu\(_2\)(OH)\(_3\)Cl species. In particular, the band at 913 cm\(^{-1}\) confirmed the presence of atacamite, while the one at 863 cm\(^{-1}\) can be attributed to clinoatacamite and anatacamite. The ATR–FTIR transmittance spectra of the CHT/IS coating, before and after 6 h of accelerated corrosion, are reported as an example in Fig. 9.

On the contrary, no changes on the alloy substrates were observed in the case of the coatings with BTA and the BTA-IS mixture (Fig. 8). Prolonging the corrosion treatment up to 12 h, the coating CHT/BTA-IS was still stable and able to prevent modifications of the metal substrate, whereas in the case of the coating CHT/BTA significant alterations were detected on the alloy surface. Some representative optical images are reported in Fig. 10. Furthermore, the ATR–FTIR spectroscopy analysis of the coating after 12 h of exposition to the acid vapors showed the formation of basic copper chlorides corrosion products (Additional file 1: Figure S6).

The superior performance of the coating CHT/BTA-IS was also confirmed by ATR–FTIR spectroscopy as no changes of the chitosan structure are observed in the FTIR spectrum after exposure to the aggressive vapors for 12 h. Moreover, FTIR bands attributable to the presence of corrosion products were not detected in the spectrum of the CHT/BTA-IS coating after corrosion treatment (Fig. 11).

In order to get more information about the coatings and the metal surfaces after the corrosion test, the coated alloy disks were further investigated by SEM–EDS
analysis. The SEM micrographs (Fig. 12) of the coating CHT/BTA-IS were recorded near the area where the film was removed after the corrosion treatment. It is clearly visible that the coating was still homogeneous and crack-free even after the prolonged exposure to aggressive vapors. Moreover, the alloy surface did not show significant modifications after film removal.

On the contrary, the SEM micrographs of the CHT/BTA coating reveal both the alteration of the film and the formation of degradation products on the alloy substrate after film removal (Fig. 13). EDS analysis confirmed the presence of corrosion products and species containing chloride on the metal surface where the coating was removed after exposure to acid vapors (Fig. 13a, b).

In order to quantify the protective efficacy of the studied coatings, the image analysis has been performed to estimate the percentage of corroded surface, and the results are given in Fig. 14. Evidently, the pristine CHT coating CHT was not able to provide durable protection against corrosion, despite its very good adhesion to the bronze surface. The addition of the IS did not improve the coating protection ability, which remained essentially equal to that of the pristine CHT coating. As expected, the presence of the BTA corrosion inhibitor significantly enhanced the ability of the coating to preserve the underlying metal surface from corrosion. Actually, the presence of corrosion products at the metal/coating interface was hardly distinguishable up to 6 h of accelerated corrosion treatment. For treatments of prolonged duration, i.e. 12 h, non-negligible surface modification occurred. Interestingly, the coating containing both BTA and the IS (CHT/BTA-IS) provided the best corrosion inhibition, thus suggesting a synergistic effect of the two once used together. Overall, the time evolution of the percentage of corroded surface was similar to that of CHT/BTA. Nonetheless, it is worth noting that the amount of corroded surface was significantly lower with CHT/BTA-IS for longer accelerated corrosion treatments.

The obtained results show that the coating CHT/BTA-IS is able to protect the copper alloy surface, avoiding the formation of degradation product, even in extremely aggressive conditions. These results also suggest that HO₂CC₄MImNTf₂ plays an important role increasing the stability and the protective properties of the CHT/BTA formulation.
Conclusion

In this work, IS $\text{HO}_2\text{CC}_1\text{MImNTf}_2$ was investigated as potential novel corrosion inhibitor. Although this IS is less effective than BTA in the protection of bronze substrate, the use of $\text{HO}_2\text{CC}_1\text{MImNTf}_2$ in combination with BTA improves the protective efficacy of the CHT-based formulation, showing a synergistic effect of the two additives. This result is of particular interest for the development of more sustainable and less toxic materials for the conservation of cultural heritage, including BTA-free coatings.

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10.1186/s40494-020-00381-4.

Additional file 1. $^1$H NMR spectra of $\text{HO}_2\text{CC}_1\text{MImCl}$ (Figure S1) and $\text{HO}_2\text{CC}_1\text{MImNTf}_2$ (Figure S3). $^{13}$C NMR spectra of $\text{HO}_2\text{CC}_1\text{MImCl}$ (Figure S2) and $\text{HO}_2\text{CC}_1\text{MImNTf}_2$ (Figure S4). Thickness determination by SEM for CHT and CHT/BTA (Figure S5). ATR-FTIR spectra of CHT/BTA coating.
on bronze disk before and after 12 h of accelerated corrosion treatment (Figure S6).

Abbreviations

ATR–FTIR: Attenuated total reflectance–Fourier transform infrared; BTA: Benzotriazole; CTR: Control; D: Doublet; DGL: D-(1-Methoxycarbonylmethyl-3-methylimidazolium chloride); MS–ESI: Mass spectrometry–electron spray ionization; NMR: Nuclear magnetic resonance; NTf2: Bis(trifluoromethylsulfonyl) imide anion; OM: Optical microscopy; RH: Relative humidity; r.t.: Room temperature; s: Singlet; SEM: Scanning electron microscopy; t: Triplet; qua: Quartet; qui: Quintet; ∆: Change in a variable; H3CO2C‑CHT/IS: 1-Carboxymethyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide; CHT: Pure chitosan coating; CHT/BTA: Coating of CHT with corrosion inhibitor BTA (2.0 wt% with respect to CHT); CHT/IS: Coating of CHT with IS H3CO2C‑CHT; CHT/IS with BTA (2.0 wt% with respect to CHT); CHT/BTA/IS: Coating of CHT with BTA (2.0 wt% with respect to CHT) and HO2C‑CHT/IS: 1-Carbosyethyl-3-methylimidazolium chloride; H3CO2C‑CHT/IS with IS: Pure chitosan coating; CHT/BTA: Coating of CHT with corrosion inhibitor BTA (2.0 wt% with respect to CHT); CHT/IS: Coating of CHT with IS H3CO2C‑CHT/IS: 1-Carbosyethyl-3-methylimidazolium bis(trifluoromethylsulfonyl)imide; CHT. Pure chitosan coating, CHT/BTA: Coating of CHT with corrosion inhibitor BTA (2.0 wt% with respect to CHT); CHT/IS: Coating of CHT with IS H3CO2C‑CHT/IS: 1-Carbosyethyl-3-methylimidazolium chloride; CHT, Pure chitosan coating.

Acknowledgements

The authors thank Mr. Mario De Angiulli for his technical support during nanoscratch analyses, Dr. Cristina Riccucci and Dr. Marianna Pascucci for field emission SEM–EDS analyses, and Mrs. Luciana Cerri for AFM measurements.

Authors’ contributions

LA, ML, GDC, MPS, TF, and HSS designed research; DKSC, KNA, MGR, CS, and CG performed research, and DKSC, EN, MSL, CG, GDC, and HSS wrote the paper. All authors read and approved the final manuscript.

Funding

This research was carried out within the framework of the NANORESTART project funded by the European Union’s Horizon 2020 research and innovation program under agreement No. 646063. This study was financed in part by: the Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES)-Brazil, Finance Code 001; the Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq)-Brasil, PIBIC; and the Fundação de Amparo à Pesquisa do Estado de Rio Grande do Sul (FAPERGS)-Brazil, PROBIC.

Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Competing interests

The authors declare that they have no competing interests.

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Received: 3 February 2020 Accepted: 7 April 2020

Published online: 22 April 2020

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