Effect of Mn Doping on the Microstructure and Electrical Properties of Potassium Niobate Ceramics Using Plasma Spraying

Rui Gao 1,2, Weiling Guo 2, Hongxing Wang 1, Xuewu Li 1,* and Zhiguo Xing 2,*

1 School of Mechanical Engineering, Xi’an University of Science and Technology, Xi’an 710048, China
2 National Key Lab for Remanufacturing, Army Armored Forces Institute, Beijing 100072, China
* Correspondence: lixuewu55@xust.edu.cn (X.L.); xingzg2011@163.com (Z.X.)

Abstract: KNN piezoelectric ceramics are of great importance in the field of scientific research due to their high Curie temperature, good electrical properties, etc. The application of potassium sodium niobate $K_{0.5}Na_{0.5}NbO_3$ (KNN) is strictly limited due to the volatility of $Na^+$ and $K^+$ in KNN and its leakage current. In order to investigate the effect of Mn doping on KNN-based piezoelectric ceramic coatings, KNN and KNN-0.02Mn (KNMN) coatings were successfully prepared using a plasma spraying technique. The phase structure, microscopic morphology, and electrical properties of the coatings were studied in the research. The results showed that both the KNN and KNMN coatings had chalcogenide structures. The KNN coating had an orthogonal phase structure, whereas the KNMN coating had a tetragonal phase structure. Compared with the KNN coating, the microhardness of the KNMN coating was improved through doping with Mn ions. The doping of Mn ions could replace both the A and B sites in the KNN piezoelectric ceramics, further reducing the oxygen vacancies and leakage currents in the coating, and its dielectric properties were improved.

Keywords: plasma spraying; doping; dielectric properties; ferroelectric properties; KNMN coatings

1. Introduction

Piezoelectric materials are important functional materials for the conversion of mechanical and electrical energy, and have a wide range of applications in various electronic components, such as sensors, actuators, ultrasonic transducers, resonators, filters, buzzers, and electronic igniters. Meanwhile, piezoelectric ceramics play an important role in scientific research due to their efficient piezoelectric effect as well as their low price and good electrical properties [1–4]. Currently, widely used piezoelectric ceramic materials are mainly comprised of lead zirconate titanate (PZT). The advantages of these piezoelectric ceramics are their excellent electrical properties and the possibility of industrial mass production. However, the Pb ions contained in the ceramics can cause a lot of harm to the environment. Thus, lead-free piezoelectric ceramics have been widely studied to replace lead-based piezoelectric ceramics [5,6]. Potassium sodium niobate (KNN) piezoelectric ceramics are currently considered as a viable alternative to PZT. Although the performance of pure KNN is inferior to that of PZT, its relatively high Curie temperature, good electrical properties, and high electromechanical coupling coefficient are the driving forces for KNN research [7–9].

However, there are still some practical problems that hinder the industrial application of KNN. The volatilization of KNN components is considered to be the reason for the low density and high defect concentration of KNN ceramics, which adversely affect their conductivity and ferroelectric performance behavior [10,11]. At the same time, leakage current plays an important role in polarization switching, as well as polarization processes, making it difficult for materials to obtain optimal piezoelectric properties and rendering them unsuitable for industrial applications [12]. Mn ions are often used as dopants in...
Pb-based materials for mechanical quality factor enhancement and dielectric loss reduction [13,14]. Huang et al. [15] used a solid-state reaction method to prepare Mn-ion-doped Pb(Mg_{0.5}W_{0.5})O_3-Pb(Zr, Ti)O_3(PMW-PZT) ceramics. The study showed that the electrical properties of the ceramics were effectively improved. Peng et al. [16] adopted the sol–gel method to prepare Mn-ion-doped PZT piezoelectric ceramic films. The results showed that 1 mol% Mn could achieve sender doping, which was beneficial to the domain movement and further enhanced the ferroelectric properties of the PZT films. In addition, Mn plays an important role in lead-free piezoelectric ceramics, such as BaTiO_3 and KNbO_3, by enhancing the densities, dielectric losses, and electrical properties of ceramics [17,18]. Current research on KNN-based piezoelectric ceramics focuses on bulk ceramics, single crystals, and thin films. Mn can reduce the leakage current density of potassium sodium niobate, which has little effect on the dielectric properties of doped KNN ceramics [19]. Wang et al. [20] used chemical solution deposition to prepare KNN thin films with different Mn doping contents. The results showed that the doping of Mn ions reduced the KNN leakage current density and the electrical properties were effectively improved, which, in turn, enhanced the ferroelectric properties of the films [21]. Shweta et al. [22] prepared K_xNa_{1-x}NbO_3 thin films with a pulsed laser deposition technique. The effect of the deposition parameters on the structure and morphology of KNN thin films was investigated. The results showed that the KNN films had a dielectric constant of approximately 531 in the high-frequency region (>500 kHz) and a residual polarization of 8.63 µC cm^{-2}. Although a solid-phase sintering method, hot-pressure sintering method, and laser pulse deposition technology have been applied in current KNN-based piezoelectric ceramics preparation techniques, and extensive research on Mn in KNN-based systems has been conducted, the effect of Mn on the properties of KNN-based piezoelectric ceramics prepared using a thermal spray technique is still unclear.

Plasma spraying is a thermal spraying method in which a piezoelectric ceramic powder is fed into a plasma jet, heated and accelerated in it, and sprayed onto different substrates to form coatings [23,24]. Due to its characteristics of a high-ion flame flow temperature, fast spraying particle velocity, and controllable thickness, plasma spraying can ensure the density of the coating and reduce its porosity under the premise of the full melting of the sprayed powder, thus, achieving a wider application prospect [25,26]. Chen et al. [27] prepared KNN-based piezoelectric ceramic coatings with a single chalcogenide phase and a dense morphology with a thermal spraying process. The structure, morphology, and properties of the coatings deposited under different conditions were investigated, and their excellent piezoelectric properties were demonstrated. Chen et al. [28] obtained a large effective piezoelectric coefficient (d_{33}) of 125 µm/V by thermal spraying KNN-based ceramic coatings onto steel substrates with a thermal spray process for the first time after the introduction of a nickel- and yttria-stabilized zirconia interlayer. Zhou et al. [29] prepared KNN-based piezoelectric ceramic coatings using supersonic plasma spraying equipment. The research results showed that the coatings had a good quality, as well as good electrical properties. Song et al. [30] prepared Li-ion-doped KNLN-based piezoelectric ceramic coatings using supersonic plasma spraying technology. The coatings had good electrical properties, with a dielectric constant of 375 and a stable dielectric loss of 0.01, and Li ion doping could successfully build a polycrystalline phase interface with the coexistence of an O–T phase boundary. Therefore, the plasma spraying technique provides a new prospect for the preparation of KNN-based piezoelectric ceramics.

The current research on the preparation of KNN-based piezoelectric ceramics using thermal spraying technology is relatively rare. In this paper, MnCO_3-doped KNN piezoelectric ceramics were prepared by using plasma spraying technology. The effects of Mn ion doping on the microstructure, mechanical properties, and electrical properties of KNN piezoelectric ceramics were investigated to reveal the mechanism of the effect for Mn ion doping on coating properties. In addition, the effect of oxygen vacancies on the ferroelectric properties was discussed in detail.
2. Experimental Procedure

2.1. KNN Powder and Coating Preparation

K$_{0.50}$Na$_{0.50}$NbO$_3$ (KNN) and K$_{0.50}$Na$_{0.50}$NbO$_3$-0.02Mn (KNMN) powders were adopted as raw materials (Quanzhou Qijin New Material Technology Limited Company, Quanzhou, China). Based on previous research on KNN piezoelectric ceramics, 10 mol% more K and Na was added considering the volatilization of alkaline metal ions during the thermal spraying process. The plasma spraying equipment had the advantages of a fast spraying speed and high spraying temperature, and its schematic diagram is shown in Figure 1.

![Schematic diagram of the plasma spraying process.](image1)

Since plasma spraying technology is suitable for the preparation of ceramic coatings due to the high temperature and speed of the jet flame center, which can melt many refractory materials, the spraying process requires a certain size and fluidity of powder, and this, in turn, affects the quality of the ceramic coating. A powder particle size that is too small can form an agglomeration in the process of powder charging or be directly melted in the process of spraying due to the high temperature of the jet flame center. As a result, the particles cannot be deposited in the substrate. Therefore, the powder had to be atomized using granulation, sieving, etc., so that the appropriate KNN and KNMN powder particle sizes could be achieved, as shown in Figure 2. It can be seen that the particle sizes of the two powders were mainly concentrated at approximately 30–80µm, where $d_{(0.1)} = 0.722$, $d_{(0.5)} = 40.059$, and $d_{(0.9)} = 77.972$; the preparation of the powder was fully suitable for the plasma spraying system.

![KNN and KNMN powder particle size.](image2)
The sprayed substrate was selected from the 45th steel with a processing size of 25 mm × 20 mm × 5 mm. Before spraying, the surface of the substrate specimen was ultrasonically cleaned with acetone and ethanol to remove oil contamination, and then sandblasted with aluminum oxide to remove oxides and other impurities to increase the roughness of the substrate surface. Next, Ni-based alloy compounds were sprayed onto the substrate surface to increase the bond strength between the substrate and the coating. Prior to spraying, the substrate surface was prepped by plasma spraying it for preheating to a temperature of 90–110 °C with the preheating time set to 15 s. Plasma spraying equipment was applied in the research and the spraying process parameters are listed in Table 1, displaying a spraying thickness of 250 µm. To enhance the electrical properties of the coating, it was heated at 600 °C in a KSL-1100X-S chamber furnace for approximately 30 min.

Table 1. Spraying process parameters.

<table>
<thead>
<tr>
<th>Process Parameters</th>
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<tr>
<td>Ar gas flow (m³/h)</td>
<td>50</td>
</tr>
<tr>
<td>Spraying voltage (V)</td>
<td>96</td>
</tr>
<tr>
<td>H₂ gas flow (m³/h)</td>
<td>8</td>
</tr>
<tr>
<td>Spraying current (A)</td>
<td>400</td>
</tr>
<tr>
<td>Spraying distance (mm)</td>
<td>100</td>
</tr>
<tr>
<td>Powder feeding speed (g/min)</td>
<td>20</td>
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2.2. Powder and Coating Performance Testing

The particle size of the powders was analyzed with a laser particle sizer (Mastersizer 2000 model); the 3D morphology of the coatings was obtained with a 3D morphological profiler (Nanovea ST400, Nanovea, Irvine, CA, USA) at a magnification of 200 times; the morphology of the powder, as well as the cross-sectional morphology of the coating, was obtained using a field emission scanning electron microscope (SEM; ZEISS Gemini, Oberkochen, Germany). Chemical elements were analyzed using energy-dispersive spectroscopy (EDS, ZEISS Gemini, Oberkochen, Germany); the phase structure of the powders and coatings was analyzed with an X-ray diffractometer (D/MAX-RB; Tokyo, Japan) at a wavelength of λ = 0.15406 nm, a Cu target, a scanning speed of 4°/min, and a scanning angle of 20°–60°. The microhardness of the two coatings was tested using a microhardness tester (KELITI000ZB, Sichuan, China). The bond strength of the coatings was tested using an MTS fatigue tester. Upper and lower electrodes were prepared to test the electrical properties of the KNN and KNMN coatings. Bottom electrodes 3 mm in diameter were prepared using an ion sputtering method. The specimens were polarized using high-temperature polarization equipment (KSL-1100X, Hefei, China). The dielectric constant and dielectric loss were tested at room temperature using an impedance analyzer (Agilent 4294A, NANOVEA, Irvine, CA, USA) at a frequency of 1 kHz. In addition, the ferroelectric properties of the coatings were tested using a ferroelectric analyzer (TF Analyzer 2000E, Beijing, China). An X-ray electron spectrometer (XPS; Thermo escalab 250XI, USA) was used to analyze the elemental valence states of the coatings. In addition, the following applied: excitation source: Al Kα (hv = 1486.6 eV); electron emission angle: 55°; test area: 650 µm × 650 µm; Ar+ energy: 2000 eV; angle: 50°; analysis pressure: 10⁻¹⁰ mba. To reduce the error, we referred to the Fermi edge (Eᶠᴮ) of the critical carbon used in Liviu’s work [31] for the calibration. The relationship between the work function of the sample (Φˢᴬ) and the Fermi edge (Eᶠᴮ) was Eᶠᴮ = 289.58 eV-Φˢᴬ. Since the work function of the piezoelectric ceramic was Φˢᴬ = 4.8 eV [31], Eᶠᴮ = 284.78 eV was used as the reference value for the BE.

3. Results and Discussion

3.1. Organizational Structure and Phase Structure

The roughness of the coating surface was one of the factors influencing the appearance of the coating, as well as its performance in use. Figure 3 represents the three-dimensional
morphology of the KNN and KNMN coatings. As shown in Figure 3, the surface of the sprayed substrate specimen was uneven and had a certain degree of roughness. In the whole spraying process, the isoelectronic arc heat between the KNN powder and the KNMN powder was exchanged through the spray gun, thus, turning the powder into a molten or semimolten state. In the plasma spraying, the powder’s movement was accelerated in the flame. The KNN and KNMN powders could be melted due to the ultra-high temperature of the flame, so that the spraying process could be completed in a short period of time. The three-dimensional shape of the coating and the flatness of the surface were closely related to the molten powder’s distribution on the substrate surface.

![Figure 3](image_url)

**Figure 3.** (a) Three-dimensional morphology of KNN coating. (b) Three-dimensional morphology of KNMN coating.

As can be seen from Figure 3, the KNMN coating had a better spraying effect and a flatter coating surface. The difference between the highest point and the lowest point of the vertical height was only 29 µm, which was because the Mn ions were a sintering aid that could melt the KNN powder at a high temperature, and, then, could better accumulate it onto the surface of the substrate.

Figure 4a shows the cross-sectional morphology of the KNMN and KNN coatings. It is clearly visible that the thickness of both coatings was approximately 220 µm. No significant cracks existed between the two coatings and the substrate due to the introduction of the transition layer, which was an improvement in terms of the process. The porosity of the coatings was measured using image-pro software. The porosity of the KNN and KNMN coatings was obtained as 5.8% and 5.2%, respectively. With better quality and lower porosity, the KNMN coatings promoted the melting of the powder due to the doping of Mn, which was consistent with the results of the tests of the 3D morphology. Figure 4c represents the EDS of the KNMN-coated sample, from which it can be seen that the main elements were K, Na, Nb, O, and Mn. The distribution of the elements was relatively uniform and corresponded exactly to the coating structure of the sample. Table 2 indicates the content of elements in the KNMN coating, which showed that the elements in the coating and the powder remained consistent. The Mn ions went into the KNN coating. The volatilization of the elements came from the high temperature during the plasma spraying process, and C from the impurities brought on by the spraying process.

<table>
<thead>
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<th>Table 2. Element contents in the coatings.</th>
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<tr>
<td><strong>Element</strong></td>
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<td>wt%</td>
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<td>at%</td>
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was observed in the XRD patterns [32]. Figure 7b represents the extended XRD patterns of the KNMN coating. Although the orthorhombic structure was the dominant phase at room temperature, it can be seen from Figure 5 that the distribution of the K, Na, Nb, O, and Mn elements was relatively uniform. The figures showed that the distribution area of the Nb and O elements was relatively large, which was consistent with the properties of the original powder, where the content of the K and Na elements was low because of the high level of volatilization of alkaline metal ions caused by high temperatures during the plasma spraying process.

Figures 5 and 6 represent the TEM/EDX surface scan element distribution and X-ray energy spectrum analysis of the KNMN coating, and the insets indicate the atomic percentages of each element of the KNMN coating; it can be seen from Figure 5 that the distribution of the K, Na, Nb, O, and Mn elements was relatively uniform. The figures showed that the distribution area of the Nb and O elements was relatively large, which was consistent with the properties of the original powder, where the content of the K and Na elements was low because of the high level of volatilization of alkaline metal ions caused by high temperatures during the plasma spraying process.

Figure 5. TEM/EDX surface scan elements of KNMN coating.

Figure 4. (a) Cross-sectional morphology of KNN coating. (b) Cross-sectional morphology of KNMN coating. (c) EDS surface scan of KNN coating.

Figure 7a represents the XRD patterns of the KNN and KNMN powders. It was concluded that the KNN and KNMN powders both had a single chalcogenide structure. The Mn ions completely entered the KNN lattice because no formation of the second phase was observed in the XRD patterns [32]. Figure 7b represents the extended XRD patterns of the KNN and KNMN powders in the range of 2θ from 44° to 47°. Both powders had an orthoclase structure as the front peak was higher than the back peak at approximately 45°.

Figure 8a represents the XRD patterns of the KNN and KNMN coatings. Both coatings showed a single chalcogenide structure. The KNMN coating exhibited a preferential orientation of 100. Figure 8b represents the extended XRD patterns of the KNN and KNMN coatings in the 2θ range of 44°–47°. It could be concluded that the KNN coating showed an orthorhombic phase structure, while the KNMN coating displayed a tetragonal phase structure. Although the orthorhombic structure was the dominant phase at room temperature, with the doping of the Mn ions it changed to a structure dominated by a tetragonal phase structure. At the same time, the tetragonal orthogonal phase transition (T0,T1) shifted to a lower temperature. The single chalcogenide phase also indicated that Mn could enter the KNN lattice by replacing the A site. It can be seen that the splitting
phenomenon disappeared in the ceramics and that the diffraction peaks shifted toward smaller angles after the addition of a quantitative amount of Mn ions. Mn could replace both the A and B sites in KNN, thus, causing a distortion of the lattice structure and defects in the oxygen ions. In addition, the peak intensity of the KNMN coating was lower than that of the KNN coating, but the doping of the Mn ions enabled the KNMN coating to maintain a better crystal structure.

![Figure 6. X-ray energy spectrum analysis of KNMN coating.](image1)

![Figure 7. (a) XRD patterns of KNN and KNMN powders. (b) 44°–47° XRD extended patterns.](image2)
which increased the coating density and, thus, increased the microhardness, indicating that the KNMN coating was of better quality. The microhardness of the coating reflects the quality of the coating. As shown in Figure 9b, the microhardness of the transition layer, which was due to the fact that the material of the transition layer was NiCrCoAlY, so the microhardness of the coating was greater. The microhardness of the coating was greater than that of the KNN coating because the doping of manganese ions during the spraying process promoted the melting of the ceramic powder, which increased the coating density and, thus, increased the microhardness, indicating that the KNMN coating was of better quality.

3.2. Mechanical Properties

The microhardness of the KNN and KNMN coatings is shown in Figure 9. From Figure 9a, it can be seen that the microhardness of the coatings tended to increase and then decrease from the coating to the substrate direction. The highest microhardness was in the transition layer, which was due to the fact that the material of the transition layer was NiCrCoAlY, so the microhardness of the coating was greater. The microhardness of the coating reflects the quality of the coating. As shown in Figure 9b, the microhardness of the KNMN coating was greater than that of the KNN coating because the doping of manganese ions during the spraying process promoted the melting of the ceramic powder, which increased the coating density and, thus, increased the microhardness, indicating that the KNMN coating was of better quality.

Figure 10 shows the bond strength of the KNN and KNMN coatings. The average values of the KNN and KNMN coating strengths were 23.8 MPa and 29.8 MPa, respectively, which could basically meet the requirements of actual working conditions. In terms of the fracture site of the specimens, the fracture locations were both located between the coating and the substrate, which further indicated that the coating was relatively dense, with better mechanical properties, and also proved that the bonding between the coating and the substrate was mainly based on mechanical bonding. The bonding strength of KNMN was
better because of the introduction of Mn ions in the KNN coating, which promoted the melting of the powder during the spraying process and increased the coating density, thus, increasing the bonding strength of the coating.

![Bond Strength of KNN and KNMN Coatings](image1)

**Figure 10.** Bond strength of KNN and KNMN coatings.

### 3.3. Electrical Properties

#### 3.3.1. Dielectric Properties

Figure 11 shows the frequency dependence of the dielectric constant and loss angle tangent of the KNN and KNMN coatings. As shown in Figure 11, the dielectric constant and dielectric loss tended to decrease and become stabilized with an increase in frequency, while the dielectric constant of the KNN coating was set at approximately 900 and the dielectric loss at approximately 0.08. In contrast, the dielectric constant of the KNMN coating could be stabilized at approximately 1700, which was significantly higher than that of the pure KNN coating, and the dielectric loss value was stabilized at approximately 0.01, which was apparently lower than that of the pure KNN coating. This indicated that the doping of the Mn ions could enhance the coating’s dielectric constant and reduce the dielectric loss.

![Dielectric Spectrum of KNN and KNMN Coatings](image2)

**Figure 11.** Dielectric spectrum of KNN and KNMN coatings.
3.3.2. Ferroelectric Properties

To investigate the ferroelectric properties of the KNN and KNMN samples, the polarization versus electric field (P–E) hysteresis loops were measured at 100 Hz, as shown in Figure 12. Both coatings still maintained intact hysteresis loops without an electric field breakdown. The residual polarization and coercivity fields of the pure KNN coating were 1.17 \( \mu \text{C/cm}^2 \) and 68 kV/cm, respectively, while the residual polarization and coercivity fields of the KNMN coating were 2.71 \( \mu \text{C/cm}^2 \) and 141 kV/cm, respectively. The residual polarization of the KNMN coating increased and the coercivity field decreased. The doping of the Mn ions could enhance its ferroelectric properties, which was due to the fact that the doping of the Mn ions could reduce the oxygen vacancies in the coating and reduce the leakage current of the coating, thus, enhancing its ferroelectric properties [33,34]. The crystallinity of the coating may have reduced the electrical properties of the coating, but the addition of the Mn ions compensated for the effect of crystallinity on the electrical properties and maintained a better crystal structure, so the electrical properties were, subsequently, improved.

![Figure 12. Electric hysteresis lines of KNN and KNMN coatings.](image)

3.3.3. XPS Results

In order to investigate whether Mn entered the lattice of the KNN chalcogenide structure and to study the valence states of the elements, the binding energies of K, Na, Nb, O, and Mn of the KNMN coating and the valence states of each element were studied using the XPS technique, and the XPS profiles of the individual elements are shown in Figure 13. Since most of the Mn was eventually transformed into a higher valence state and because its radius was smaller than the interaction between K, Na, and Nb, the interaction between Mn and O was stronger than that between K, Nb, Na, and O. Therefore, the electron cloud density around oxygen decreased, leading to an increase in the K, Na, and Nb binding energy of the Mn-doped KNN films [16]. According to the results of the XPS fitting, peaks were found at 640.35 eV, 641.59 eV, and 642.7 eV and 651.95 eV, 653.11 eV, and 654.14 eV, as shown in Figure 13, which corresponded to Mn\(^{2+} \), Mn\(^{3+} \), and Mn\(^{4+} \), respectively [21]. The peak area indicated the content of a specific ion, but, with advantages, the exact area of the peak could not be accurately calculated and the height of the area was set according to the maximum value of the half-peak width. In contrast, the area of 640.35 eV was smaller than that of 641.59 eV and 642.7 eV, and so, it could be concluded that the first-introduced Mn\(^{2+} \) ions were mainly oxidized to higher valence ions at high temperature.
Figure 13. XPS pattern analysis of KNMN coating: (a) XPS full spectrum; (b) Cl spectrum; (c–g) K, Na, Nb, O, and Mn XPS mapping.

The presence of oxygen vacancies in the coating was due to the volatilization of alkaline ions at the A site, i.e., Na and K elements, while, at the B-site, Nb$^{5+}$ ions were reduced to Nb$^{4+}$ at high temperature, which generated oxygen vacancies, and the oxidation of oxygen vacancies generated vacancies of the main carriers; the oxidation reaction was as follows [30]:

$$ V + \frac{1}{2}O_2 \rightarrow O_O + 2h^+ $$

(1)

It has been reported that the addition of Mn to KNN piezoelectric ceramics can replace both the A site and B site in sodium potassium niobate. When Mn$^{2+}$ replaces the A site in sodium potassium niobate, the addition of Mn ions can further promote the consumption
of cavities and reduce the volatilization of alkaline elements in the A site, further reducing the leakage current as well as oxygen vacancies in the coating [30].

\[ Mn^{2+} + h^+ \rightarrow Mn^{3+} \]  
(2)

In addition [21,35], the occupation of \( Mn^{2+} \) on the B site of the potassium sodium niobate coating can lead to more oxygen vacancies, while \( Mn^{3+} \) can be oxidized to a higher valence state through absorbing vacancies (Equations (2) and (3)). The oxidation of \( Mn^{2+} \) can effectively reduce the vacancies, allowing for the conversion of oxygen vacancies to lattice oxygen, which, in turn, leads to a decrease in the leakage current density, as well as the concentration of oxygen vacancies [30].

\[ Mn^{3+} + h^+ \rightarrow Mn^{4+} \]  
(3)

The three valence states of \( Mn \) ions also illustrated the idea that the doping of Mn ions could replace both the A and B sites in sodium potassium niobate. The oxygen vacancies of the coating mainly come from the volatilization of alkaline elements at the A site of sodium potassium niobate and the replacement of sodium potassium niobate with \( Mn^{2+} \) B sites, while the presence of Mn ions can reduce the consumption of vacancies and oxygen vacancies and the leakage current of the coating, thus, further improving the electrical properties of the coating.

4. Conclusions

The KNN and KNMN coatings were prepared using plasma spraying technology in the research, and the properties of the coatings were studied. The main conclusions were as follows: (1) The KNN and KNMN coatings had an orthoclase structure; the KNN coatings had an orthorhombic phase and the KNMN coatings had a tetragonal phase structure. (2) The doping of the Mn ions could effectively promote the melting and the quality of the coating, and improved its mechanical properties. (3) The KNMN coating had better dielectric and ferroelectric properties; the dielectric constant and dielectric loss of the KNMN coating could be stabilized at 1700 and 0.01, respectively; the dielectric constant and dielectric loss of the KNN coating were stabilized at 900 and 0.08, respectively.

**Author Contributions:** R.G.: writing—original draft; investigation, and data curation. X.L.: supervision. W.G.: review and editing. Z.X.: review and editing. H.W.: editing. All authors have read and agreed to the published version of the manuscript.

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**Conflicts of Interest:** The authors declare no conflict of interest.

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