Development of a wearable infrared shield based on a polyurethane–antimony tin oxide composite fiber

Abstract
Here, we investigate a wearable-based IR and thermal stealth structure that effectively blocks IR and thermal radiation from a human body or device using a polyurethane–antimony tin oxide (PU–ATO) composite fiber. The aging time of the ATO sol prepared by a sol–gel method, and the concentration of ATO with respect to that of the PU matrix were optimized to prepare PU–ATO composite fibers that simultaneously have an appropriate mechanical strength (strength of ~4 MPa and strain of ~340%) and IR- and thermal radiation-shielding properties with ~98% IR light, as determined by Fourier transform IR spectroscopic studies. The fabricated PU–ATO composite fiber showed stable IR- and thermal radiation-shielding properties even when exposed to ten cycles of repeated temperature changes of −20°C and +80°C and long-term temperature changes for 30 days. In addition, the surface of the PU–ATO composite fiber was rendered hydrophobic to prevent the distortion of the IR and thermal radiation due to the wetting of the PU–ATO composite fiber with absorbed water. The PU–ATO composite fiber-based textile proposed herein can be applied in wearable IR- and thermal radiation-shielding technologies to shield IR signals generated by objects of diverse and complex shapes.

Introduction
Stealth technology refers to a technology that renders objects unidentifiable by reducing or distorting the objects’ external signals (radar, IR, acoustic, and visual signals) used to identify objects1–10. In particular, IR stealth is a technology that makes IR radiation emitted by an object as similar as possible to ambient background radiation such that an IR detection sensor fails to distinguish the target object. Although all objects with a temperature above absolute zero emit IR (0.78–1000 μm), most IR signals are scattered and absorbed by microparticles, water vapor, or carbon dioxide molecules in the air, and only near-IR (NIR; 0.78–2.5 μm), mid-IR (MIR; 3–5 μm), and far-IR (FIR; 8–14 μm) signals have excellent air permeability. The wavelengths of these IR signals also overlap with the wavelength range of thermal radiation (0.1–100 μm). Therefore, in accordance with the Stefan–Boltzmann law, which states that the intensity of the IR radiation is proportional to the fourth power of the object’s surface temperature, IR light irradiated on an object can relay thermal energy to that object. Therefore, the most effective stealth method for avoiding IR detection and tracking is to bring the detectable IR signals of the object to a level as similar as possible to that of the ambient background radiation by reducing the surface temperature of the IR-emitting object or by an IR-shielding material5,9,10. To achieve this, either the object should be wrapped with an insulating material to prevent heat from spreading to the surface of the object, or the
intensity of the emitted IR radiation should be reduced by coating the object with a material that can shield IR signals.

Materials used in IR-shielding coatings include metal powders, conductive polymers, organic/inorganic composites, and semiconductors. Metal powders have high visible and IR light reflectance owing to their low emissivity, but they inevitably suffer from a key drawback: their IR-shielding property deteriorates because of the facile oxidation of the metal under long-term exposure to air. In addition, because metals are heavy, they easily increase the total weight of the object to which they are applied, which is often disadvantageous. Conductive polymers and organic/inorganic composites could be shaped into various shapes/forms; as they are flexible, they can also be used as wearable materials. However, they suffer from some drawbacks: they have poor IR-shielding capabilities and mechanical properties, and the binder preparation process is also complicated. On the other hand, semiconductors with a wide bandgap are more stable than polymers and composites in the ambient atmosphere, and have high reflectivity in the IR region. They are therefore frequently used as IR-shielding materials. Moreover, such semiconductors can also be used as wearable materials when blended with an appropriate polymer. Recently, studies on the IR-shielding properties of titanium oxide, zinc oxide, indium tin oxide, tin oxide (SnO2), and antimony tin oxide (ATO) have been reported. Among these, ATO is one of the typical semiconductors with visible-light transmittance and IR-shielding properties; owing to its affordable price, ATO is widely used in IR- and thermal radiation-shielding coatings. Currently, ATO nanoparticles are synthesized by a variety of techniques, such as hydrothermal, co precipitation, combustion, sol–gel, and direct current arc plasma jet methods, and in general, the nanoparticles are mixed with a binder to form films that can be used for IR and thermal radiation shielding. However, there is a lack of studies on the fabrication of ATO nanoparticles in a fiber form, instead of the typical film form, for application in wearable IR- and thermal radiation-shielding clothing. Moreover, there are reports on using a binder to coat ATO nanoparticles on a polymer-based fiber or mixing ATO nanoparticles with a polymer matrix. However, when a binder is used to coat ATO nanoparticles on the fiber, the coating might be damaged under friction. When ATO nanoparticles are mixed with a polymer matrix, the mechanical properties of the fiber may degrade. Accordingly, for application in wearable IR- and thermal radiation-shielding clothing, it is necessary to develop a shielding material that can effectively increase the shielding efficiency while maintaining the mechanical strength of the fiber without significantly increasing the fiber weight.

Here, we investigated the IR- and thermal radiation-shielding properties of fibers composed of a polyurethane (PU)–ATO composite. An optimal ATO sol was obtained by controlling the concentration of the ATO precursors, and aging time of the sol prepared by a sol–gel method; this sol was mixed with preset amounts of the PU solution to obtain PU–ATO composite suspensions with different compositions. A wet-spinning technique was used to fabricate PU–ATO composite fibers from composite suspensions with four different ATO concentrations. Then, the IR- and thermal radiation-shielding efficiency and mechanism of the prepared fibers were evaluated. We also determined whether the PU–ATO composite fibers could be used as a wearable material by testing their typical mechanical and optical properties according to changes in the aging time and the ATO content. Moreover, the surface of the PU–ATO composite fiber was chemically modified to render it hydrophobic, in order to resolve the problem of the distortion of the IR- and thermal radiation-shielding function when the shielding fiber is exposed to atmospheric water. Furthermore, the functional reliability of the fabricated PU–ATO composite fiber was evaluated by repeatedly testing it under applied IR and thermal radiation.

Materials and methods

Fabrication of the PU–ATO composite fiber

A 16.7 wt% PU solution was prepared by dissolving PU granules (Avention) in dimethylformamide (DMF, Sigma-Aldrich) and tetrahydrofuran (THF, Sigma-Aldrich) cosolvent (3:2 ratio). The prepared PU solution was stirred for 6 h. An ATO sol was prepared by mixing 0.5, 1.5, 3.0, and 4.0 mmol of tin(II) chloride dihydrate (SnCl2·2H2O, Sigma-Aldrich) and 0.05, 0.15, 0.3, and 0.4 mmol of antimony(III) chloride (SbCl3, Sigma-Aldrich), respectively, in 1.4 g of DMF–THF cosolvent (3:2 ratio). Each ATO sol preparation was stirred for 6 h in a closed container to homogenously mix the constituents and then aged in air for 1, 2, or 3 days. The PU–ATO composite suspensions were prepared by adding one of the prepared ATO sols to 5 g of the PU solution. Then, the composite suspension was loaded into a 10-mL disposable syringe installed in a syringe pump (Invisible Co.). Thereafter, the composite suspension was continuously injected into a water coagulation bath at a rate of 70 mL/min through an 18-gauge stainless steel syringe needle. To ensure complete removal of the solvent in the extracted PU–ATO composite fibers, the fibers were kept in the coagulation bath for 20 min. Then, the fibers were removed from the bath and dried at room temperature (20 °C) for 12 h. After drying, the PU–ATO composite fibers were immersed in a 5 mM solution of HDF-PA (Apollo Scientific) in isopropyl alcohol (JT Baker) for 30 min. Then, the PU–ATO composite fiber with HDP-PA
was dried at room temperature (20 °C) for 6 h. Finally, the dried PU–ATO composite fiber was thermally annealed for 1 h at 150 °C using a forced convection drying oven (OF-02GW, Jeio Tech).

Analysis of the optical, electrical, and mechanical properties of the PU–ATO composite fiber

The average size of the ATO sol synthesized by the sol–gel method was analyzed by DLS (Nanotrac Wave, Microtrac), while its crystallinity was analyzed by XRD. Light transmission in the NIR region was analyzed using a UV–vis–NIR spectrophotometer (LUX-2000, Invisible Co.). The surface morphology and size of the PU–ATO composite fibers were characterized by FE-SEM (S-4800, Hitachi), while chemical mapping for the identification of the constituents of the fiber was carried out by EDS (7593-H, Horiba). The surface morphology and elemental distribution of the PU–ATO composite fiber were analyzed by AFM. The water wettability of the composite fibers was assessed using a contact angle analyzer (Phoenix 300, SEO Co.). Thermogravimetric analysis (TGA/DSC 1, Mettler Toledo) of the PU–ATO composite fibers was performed under a nitrogen atmosphere. Mechanical properties, such as the tensile strength and strain, were evaluated at room temperature using a TMA (TMA7100, Hitachi). IR transmission was analyzed using an ATR–FTIR spectrometer (Tensor 27 Spectrometer, Bruker). A hot plate (MSH-20D, DAIHAN Scientific) was used as the heating source, and a Peltier module was used as the cooling source. The IR– and thermal radiation-shielding properties of the textiles were analyzed by imaging using a thermal IR camera (T-420, FLIR) after placing the textiles 2 mm from the heating and cooling source.

Results and discussion

According to Wien’s displacement law ($\lambda_m = \frac{2897.8}{T}$), if an object is assumed to be a black body, the maximum wavelength of the radiation ($\lambda_m$) of the object at room temperature (near 20 °C) is ~9.9 µm, which is located in the FIR region, whereas an object at 1000 °C emits ~2.3-µm NIR light. Therefore, to render an object indistinguishable by the IR detector sensor, it is important to make the IR and thermal radiation intensity of the object being detected as similar as possible to the IR and thermal radiation shielding due to water absorption by the fibers. In the images, a part of the finger appears to have been cutoff, owing to its invisibility.
differences in the temperatures of different parts of the body. Even when the person wears typical clothing, such as polyester or PU fabric, the IR and thermal radiation emitted from the human body passes through the fabric and becomes visible. In contrast, when a part of the human body is concealed with PU–ATO composite fiber-based clothing with IR- and thermal radiation-shielding properties, the IR and thermal radiation emitted from the human body is shielded, and the person merges with the background; accordingly, the IR camera cannot recognize the person. The PU–ATO composite fiber-based textile (PU–ATO textile) developed in the present study has superior IR- and thermal radiation-shielding properties compared with those of regular fabrics, such as polyester or PU fabric. For the fabrication of the PU–ATO composite fiber, a PU–ATO composite suspension was prepared by evenly embedding an ATO sol in a PU matrix that has elongation ability, tensile strength, and a high modulus favorable for forming a fiber; then, fibers were obtained by wet-spinning the composite suspension. The PU–ATO (3) composite fibers used to make the IR- and thermal radiation-shielding textiles were fabricated under optimal conditions, as determined by the assessment of their mechanical, chemical, and optical properties.

The attenuated total reflection–Fourier transform infrared (ATR–FTIR) spectra in Fig. 1b reveal that the IR transmittance of textiles made with polyester, PU, and PU–ATO (3) are ~40, 25, and 2%, respectively. Note that the ATR–FTIR spectroscopy was performed in the wavelength range of 7–14 µm, which includes the entire wavelength range of 7.5–13 µm detected by a thermal IR camera. The low IR transmittance of the PU–ATO (3) textile is due to the uniform dispersion of the ATO sol in the PU matrix of the fiber. In general, SnO2, a typical n-type semiconductor with visible-light transparency, modulated by controlling the Sb dopant concentration, particle size, and crystallinity.21,28 The excellent IR- and thermal radiation-shielding effects of ATO arise from the localized surface plasmon resonance generated by the Sb doping effect. Here, a plasmon refers to a quasiparticle with collective oscillations of free electrons in a conductor, such as a metal. In general, plasmons are present only on the surface of nanoparticles, and accordingly, they are also referred to as surface plasmons. When IR light is irradiated on a metal nanoparticle, the light energy is altered by the surface plasmons and accumulates on the surface of the nanoparticle. Such interaction of light with nanoparticles is referred to as surface plasmon resonance.

Plasmons generated by the collective oscillation of free electrons have a very important effect on the optical properties of the material. For example, light with a frequency lower than the plasmon frequency is reflected by the oscillating free electrons in the conduction band, whereas light with a frequency higher than the plasmon frequency is transmitted. In other words, the frequency at which IR radiation can be reflected by plasmon vibration is proportional to the plasmon frequency ($\omega_p$) in accordance with Eq. (1):

$$\omega_{sp} = \frac{\omega_p^2}{1 + \varepsilon_m}$$

where $\omega_p$ is the surface plasmon frequency, and $\varepsilon_m$ is the dielectric constant of the surrounding medium. The plasmon frequency ($\omega_p$) is proportional to the square root of the free electron density ($N$) (Eq. (2)):

$$\omega_p = \sqrt{\frac{Ne^2}{m_e\varepsilon_{\infty}\varepsilon_0}}$$

where $e$ is the electron charge, $m_e$ is the effective mass of an electron, $\varepsilon_{\infty}$ is the vacuum permittivity, and $\varepsilon_0$ is the optical dielectric constant.21 Sb may exist in two different oxidation states in a solid solution of SnO2. Furthermore, the ionic radius of Sn$^{4+}$ (0.83 Å) is slightly larger than that of Sb$^{5+}$ (0.74 Å) and slightly smaller than that of Sb$^{3+}$ (0.93 Å).29 At a low doping level, the n-type dopant Sb$^{5+}$ is dominant, whereas the p-type dopant Sb$^{3+}$ is dominant at a high doping level. The doping level at which Sb$^{5+}$ transforms to Sb$^{3+}$ is known to be ~15%30–32. According to Eq. (2), ATO, which is SnO2 doped with n-type dopant Sb$^{5+}$, has an increased free electron density ($N$) compared with undoped SnO2, and its plasmon frequency ($\omega_p$) is also increased. Moreover, radiation with a frequency lower than the electron plasmon frequency ($\omega_e$) is completely reflected without passing through the material. Therefore, the IR reflectivity ratio increases because of free electrons generated by Sb$^{5+}$ ions in ATO. By employing the PU–ATO composite fiber-based textile with ATO sol evenly embedded in the PU matrix, the IR radiation emitted by an object could be effectively reflected. As shown by the inset image in Fig. 1c, the flexible PU–ATO composite woven into a textile has a mesh structure, which is more advantageous than a more densely woven conventional polyester or PU textile in terms of IR- and thermal radiation-shielding properties, as well as air permeability. In addition, because the hydrophilic surface of the ATO sol generated by the sol–gel reaction can be chemically modified to render it hydrophobic, IR distortion that can occur when the textile becomes wet can be prevented. The flexibility, mesh structure, and hydrophobicity of the PU–ATO composite
textile demonstrate that it is suitable for use as a wearable IR shield.

To increase the IR-shielding rate of the PU–ATO composite fiber, the ATO sol content in the PU matrix of the fiber was increased. As the content of the ATO constituting the PU–ATO composite solution increases, the amount of ATO sol exposed to the surface during thin-film formation increases, and the IR transmittance of the PU–ATO composite solution decreases according to the plasmon reflection properties of the ATO. Furthermore, to enhance the IR-shielding rate of the PU–ATO composite fiber, the crystallinity of the ATO was increased by increasing the aging time of the ATO sol. By increasing the crystallinity of the ATO sol, the semiconducting properties are further expressed, and the plasmon reflection effect can be increased by the collective vibration of free electrons.

To confirm the IR- and thermal radiation-shielding effect, a part of the human body was concealed with the PU–ATO (3) composite textile, and the IR and thermal radiation emitted by the human body was captured with the IR camera. The image shown on the left side of Fig. 1d is a visible-light image of a person wearing a white cotton T-shirt partially stitched with polyester, PU, and PU–ATO (3) composite-based textiles. The size of each textile is ~50 × 50 mm². The image shown on the right side of Fig. 1d is the IR image of the same person wearing the t-shirt. The image shows the appearance of various IR intensities throughout the entire T-shirt that result from the body heat from the person. Among the three areas where the textiles (polyester, PU, and PU–ATO (3)) were applied, the area where the PU–ATO (3) textile was applied shows a low IR intensity, appearing as if there is a hole in the chest. Note that when measured with the IR camera, the temperature in the chest area was found to be ~31–35 °C, whereas the temperatures measured from the area in which the three fabrics were applied were ~33 °C (polyester), ~28 °C (PU), and ~24 °C (PU–ATO (3)). The ambient temperature around the person was ~26 °C, which is comparable with the temperature measured from the PU–ATO (3) composite fiber-based textile. Based on these findings, it was confirmed that the PU–ATO (3) composite fiber-based textile could effectively shield IR and thermal radiation generated by the human body. The image shown on the left side of Fig. 1e is a visible-light image of an index finger with the middle part wrapped with the PU–ATO (3) composite fiber-based textile. The width and length of the white PU–ATO (3) textile wrapped around the index finger are ~6 and ~15 mm, respectively. The image shown on the right side of Fig. 1e is the IR image of the hand with the PU–ATO (3) textile wrapped around the index finger. In contrast to the hand-emitting IR with varying intensities due to body heat, the part wrapped with the PU–ATO (3) textile shows an IR intensity that is comparable with that of the surrounding background; it appears as if a part of the finger has been cutoff. Note that the temperature of the finger measured with the IR camera was ~37 °C, whereas the temperature in the area where the PU–ATO (3) textile was applied was ~22 °C; the ambient temperature around the hand was ~21 °C. These findings confirm that the PU–ATO (3) textile exhibits excellent properties with regard to its functionality as an IR and thermal radiation shield.

The IR-shielding studies reported thus far used a method of mixing a solid metal or oxide nanoparticle with a binding material to fiberize or coat the outside of the raw fiber. Conversely, this study used composite materials uniformly embedded with ATO, i.e., IR-shielding materials, inside the polymer. To our knowledge, this is the first study to use composite materials in which sol–gel ATO sol particles with high surface interfacial forces are uniformly embedded in a flexible polymer. As the concentration and aging time of the ATO sol increase, the crystallinity of the ATO increases, thus enhancing the IR shielding rate.

Figure 2a shows the schematic of the method used to prepare PU–ATO composite suspensions by mixing an equal amount of the PU solution with an ATO sol solution, which was prepared with different ATO precursor concentrations and aged for different durations. The size and crystallinity of ATO sol particles prepared by a sol–gel method can be altered by controlling the ATO precursor concentration and aging time, even under mild processing conditions (20 °C, 1 atm). Moreover, unlike in the case of mixing nanoparticles with a polymer matrix, the ATO sols can be incorporated into the PU matrix because of the hydrogen bonding interactions between the reactive functions on the ATO sol surface and hydrophilic PU polymer chains. Accordingly, there is no significant degradation in the mechanical properties of the polymer after fabrication of the composite fiber. In this study, to synthesize the ATO sol (SnO₂ sol doped with 10% Sb), the host precursor tin(II) chloride dehydrate (SnCl₂·2H₂O) and the dopant precursor SbCl₃ were mixed at a molar ratio of 10:1 with DMF/THF cosolvent (DMF:THF, ~3:2). The SnO₂ sol doped with 10% Sb used in this study is SnO₂ doped with Sb⁵⁺ at a low doping level, and the IR reflectivity ratio is higher than that of undoped SnO₂ because of free electrons generated by Sb⁵⁺ ions. Here, four different ATO precursor concentrations (0.5, 1.5, 3.0, and 4.0 mmol) and four different aging times (not aged and aged for 1, 2, and 3 days) were used to synthesize the ATO sols with controlled sol particle size and crystallinity. Note that ATO sols prepared by mixing 1.4 g of DMF/THF cosolvent (3:2 ratio) with 0.5, 1.5, 3.0, and 4.0 mmol of ATO precursor (0.5, 1.5, 3.0, and 4.0 mmol of SnCl₂·2H₂O with 0.05, 0.15, 0.3, and 0.4 mmol of SbCl₃, respectively) are referred to as ATO (1), ATO (2), ATO (3), and ATO (4). As shown, ATO sols
with different ATO concentrations aged for different durations were mixed with the same amount (5 g) of a 16.7 wt% PU solution in the DMF-THF cosolvent (3:2 ratio), and stirred evenly for at least 6 h to prepare the PU–ATO composite suspensions. To verify the optical properties of the prepared PU–ATO composite suspensions with different compositions, their transmittance in the NIR range was evaluated by ultraviolet–visible–near-infrared (UV–vis–NIR) spectroscopy. Figure 2b shows the transmittance in the NIR region of thin films fabricated using the PU–ATO composite suspensions and the ATO sol precursors (not sol aged) prepared by directly mixing the ATO precursors with the PU polymer. Note that the PU–ATO composite thin film was solidified by immersion in a water bath for 20 min, similar to when fabricating the fiber by wet-spinning; the thickness of the fabricated thin film was ~120 μm. Here, the ATO sols were prepared without aging, and only the concentration of ATO mixed with the PU polymer was changed. While the PU–ATO composite suspensions were mixed for at least 6 h, the ATO precursors in the PU matrix underwent hydrolysis and condensation reactions to form ATO sol particles. As shown in Fig. 2b, the FTIR results indicate that the IR transmittance tends to decrease rapidly with an increase in the concentration of ATO and then gradually saturates. This is because the amount of the ATO sol exposed on the film’s surface during thin-film formation increases with increasing ATO content in the composite suspension; the IR transmittance of the PU–ATO composite thin film decreases according to the ATO plasmon reflection properties (Supplementary Figs. S1, S2, Supplementary Information). Figure 2c, d shows
the IR transmittance of thin films fabricated by aging ATO sols of different concentrations for 1 or 2 days at room temperature (20 °C), without stirring and then mixing each of them with the PU solution. As shown in the figures, the tendency of a rapid decrease followed by gradual saturation of the IR transmittance with increasing ATO concentration is observed, regardless of the aging time for all PU–ATO composite films. However, Fig. 2e, which compares the IR transmittance of different thin films at a wavelength of 1,200 nm (data extracted from Fig. 2b–d), clearly reveals a gradual decrease in the IR transmittance with an increase in the aging time. On the other hand, when the IR transmittances of thin films with ATO sols aged for 2 and 3 days were compared, no distinct difference was observed (Supplementary Fig. S3, Supplementary Information).

Dynamic light scattering (DLS) was used to identify the average size of ATO sol particles according to the aging time. As shown in Fig. 2f, the size of ATO sol particles increased with increases in the ATO concentration and aging time. These results are consistent with the IR transmittance being inversely proportional to the ATO concentration, as shown in Fig. 2b–e. Moreover, as shown in Fig. 2g, the diffraction peaks of ATO sol particles appeared at (110), (101), (200), (211), (220), and (002) when 3 mmol of the ATO sol (referred to as ATO (3)) was aged between 1 and 3 days. The peaks are very similar to the diffraction peaks of SnO2, indicating that Sb was doped well in the lattice of SnO2 in the ATO sol. Moreover, each diffraction peak became sharper and more intense with an increase in the aging time from 1 to 3 days, suggesting that the crystallinity of ATO sol particles increased as the aging time increased. In general, during aging, sol particles grow in size as they randomly collide with each other. Cohesion between sol particles occurs through forces such as van der Waals forces between particles; if the aging time is too long, the primary particles grow into significantly larger particles and eventually sediment under the influence of gravity. As shown in Fig. 2f, the particles grew to a size larger than 1.5 μm by the third day, and some amount of the precipitate settled at the bottom of the container. When precipitates are present, the dispersion of the composite in the solvent is decreased, which could degrade the mechanical properties of the fabricated polymer composite fiber.

As the aging time of the ATO sol increased, the strain and strength of the fabricated PU–ATO composite fiber, which represent its mechanical properties, tended to decrease (Supplementary Fig. S4, Supplementary Information). This is because the size of the ATO sol particles increased as the aging time increased, and their interactions with polymer chains decreased partially as a consequence. In particular, the stability of the dispersion of the composite in the solvent decreased starting from an aging time of 3 days because of the precipitation of ATO sol particles. As a result, the strain and strength of the fabricated PU–ATO composite fiber decreased rapidly. Here, the composite dispersion in DMF/THF showed no precipitation when aged for 2 days, implying that there would be almost no degradation in the mechanical properties. Therefore, the samples aged for 2 days, which showed the best IR- and thermal radiation-shielding properties, were selected for the fabrication of the IR- and thermal radiation-shielding fiber and textile. Using PU–ATO composite suspensions prepared from ATO sols of four different concentrations (0.5, 1.5, 3.0, and 4.0 mmol) aged for 2 days, PU–ATO composite fibers were fabricated by a conventional wet-spinning method. The four types of PU–ATO composite suspensions were injected into a coagulation bath containing water through a syringe. Here, to fabricate the fiber by wet-spinning, it is important that the coagulation liquid be a nonsolvent for PU and ATO while being miscible with THF and DMF. Therefore, deionized (DI) water was used as the coagulation liquid. When the PU–ATO composite suspension in the fiber form is injected into DI water via a syringe, the concentration difference causes an exchange between the coagulation liquid (DI water) and the solvent of the composite (THF and DMF). As DI water penetrates the PU–ATO composite fiber, the solubility of the PU and ATO is gradually reduced, causing the solidification of the PU–ATO composite into a composite fiber. PU–ATO composite fibers fabricated in this manner have evenly embedded ATO sol particles in the PU matrix; additionally, because of the localized surface plasmon resonance of such ATO sol particles, the PU–ATO composite fibers reflect the IR light irradiated from the outside. When the PU–ATO composite thin films were annealed for 1 h at four different temperatures (25, 100, 150, and 200 °C), the IR transmission tended to decrease with an increase in the temperature (Supplementary Fig. S5A, Supplementary Information). Although the films annealed at 25 and 200 °C did not show a significant change in transmission (with up to only ~10% difference), the IR transmission of the composite fiber could be reduced further by a simple thermal annealing process. Accordingly, the fabricated composite fibers were thermally annealed for 1 h at four different temperatures (25, 100, 150, and 200 °C). However, the degradation of the PU–ATO composite fiber began at 200 °C (Figure S5B, Supplementary Information). It is believed that although heat treatment at a high temperature (≥500 °C), where sintering of ATO sol occurs, cannot be applied because of the degradation of the organic PU matrix, an increase in the annealing temperature up to 150 °C facilitated condensation of the unreacted precursor molecules present inside the ATO sol particles, causing an increase in its crystallinity.
Based on these findings, the fabricated PU–ATO composite fibers were thermally annealed at 150 °C for 1 h to maximize their IR- and thermal radiation-shielding properties.

As shown in the FE-SEM images in Fig. 3a, the diameters of the PU–ATO (1, 2, 3, and 4) composite fibers increased slightly as the ATO content increased: ~260 (PU–ATO (1)), ~300 (PU–ATO (2)), ~330 (PU–ATO (3)), and ~370 µm (PU–ATO (4)). This is because as the ATO content increases, the PU content in the fabricated PU–ATO composite fiber decreases. Accordingly, as the PU content decreases, alignment and shrinkage of the PU polymer chains occur to a lesser degree during the drying of fibers after wet-spinning, which causes the diameter of the fiber to increase slightly.

Figure 3b displays the atomic force microscopic (AFM) images showing the surface characteristics of the PU–ATO (3) composite fibers. The height mode AFM image of the PU–ATO (3) composite fiber shows its surface morphology with height variation within 1 µm. Moreover, the area of the ATO sol embedded in the PU matrix was identified by the phase-mode AFM image. In the phase-mode AFM image, the difference between high and low contrast indicates phase shifts in that area. The morphology of the imaged area is irregular, and the area accounts for particles of dozens of nanometers to 500 nm size on the surface of the PU–ATO (3) composite fiber, which is less than the size of the ATO sol particle (~1 µm). In the phase image of the PU–ATO (3) composite fiber, the area consisting of ATO with relatively high contrast is distributed locally inside the PU matrix with low contrast.

Figure 3c shows an optical image of a PU–ATO (3) composite fiber wrapped around a plastic rod with a diameter of 17 mm and cross-sectional FE-SEM image of the fiber. EDS elemental maps showing the distribution of carbon, tin, and fluorine in the fiber. Fig. 3 Surface structure, material composition, thermal stability, strength, IR transmission, and hydrophobic properties of PU–ATO composite fibers. a FE-SEM images of PU–ATO (1–4) composite fibers with four different ATO concentrations. b Height and phase-mode AFM images of a PU–ATO (3) composite fiber. c Optical image of a PU–ATO (3) composite fiber wrapped around a plastic rod with a diameter of 17 mm and cross-sectional FE-SEM image of the fiber. EDS elemental maps showing the distribution of carbon, tin, and fluorine in the fiber. d TGA, e TMA, f IR transmission between 7 and 14 µm, and g water contact angles of PU–ATO (1–4) composite fibers. h Photograph of water droplets dropped on top of pristine and hydrophobic PU–ATO (3) composite fiber-based textiles. i IR images of pristine PU–ATO (3) composite fiber-based textile partially wetted by water and hydrophobic-treated PU–ATO (3) composite fiber-based textile not wetted by water when heated on a hot plate at 40 °C for 1 min.
2H-heptadecafluorodec-1-yl)-phosphonic acid (HDF-PA) solution in isopropyl alcohol (IPA). During this process, self-assembled monolayers (SAMs) with strong chemical bonds are formed due to the condensation reaction between the phosphonic acid groups of HDF-PA molecules and hydroxyl groups on the surface of the ATO sol particle, which is a metal oxide. As seen in the EDS maps, carbon as the main component of PU, tin in ATO, and fluorine in the HDF-PA SAMs are evenly distributed in the PU–ATO (3) composite fiber.

Figure 3d shows the thermogravimetric analysis (TGA) curves of the PU–ATO composite fibers with four different ATO concentrations. The temperature range of TGA is 100–700 °C. The thermograms show section I between ~250 °C and 300 °C, where a sharp weight loss occurred, and section II between 300 °C and 400 °C, where a gradual weight loss occurred. In general, the PU polymer includes hard and soft segments. In section I, breakage of urethane bonding due to the pyrolysis of hard segments occurred. In section II, polyol decomposition due to the pyrolysis of soft segments occurred. At temperatures greater than 450 °C, only the inorganic high-melting ATO remains after the degradation of the polymer chains, and as a result, no further weight loss occurred. Consequently, the final remaining mass at the reference temperature of 700 °C in the TGA thermograms of the PU–ATO composite fiber tends to increase according to the ATO content from 19% (pristine PU) to 29% (PU–ATO (1)), 33% (PU–ATO (2)), 36% (PU–ATO (3)), and 38% (PU–ATO (4)).

To apply the fabricated composite fiber as a wearable textile, its mechanical properties must be verified. Figure 3e shows the tensile strength (maximum load) and strain (elongation) values at the breaking point of the PU–ATO composite fibers during the tensile test conducted using a thermomechanical analyzer (TMA). As the ATO content increased, both the tensile strength, 11.70 (PU–ATO (1)), 11.14 (PU–ATO (2)), 3.90 (PU–ATO (3)), and 1.82 (PU–ATO (4)) MPa, and the tensile strain, 434.9 (PU–ATO (1)), 374.9 (PU–ATO (2)), 335.7 (PU–ATO (3)), and 258.3% (PU–ATO (4)), of the fiber showed gradually decreasing trends. Consequently, as the ATO content increased, the interactions between the PU polymer chains became weaker, which caused a decline in the mechanical properties of the PU–ATO composite fiber because of the decreased crystallinity of the PU polymer matrix. The tensile strength of ~2–12 MPa and strain of ~250–470% for the PU–ATO (3) composite fiber are similar to those of the polyester- or PU-based fibers fabricated by conventional wet-, electro-, and melt-spinning methods (strength: 23–250 MPa, strain: ~11–330% for polyester, and strength: 6–60 MPa, strain: ~120–750% for PU).

Figure 3f shows the IR transmission in the IR region (wavelength range, 7–14 µm) of the pristine PU fiber and PU–ATO composite fibers with four different ATO concentrations, as measured by ATR–FTIR. The pristine PU fiber showed an IR transmission of ~25%; however, as the concentration of ATO with high IR reflectance increased, the IR transmission of the PU–ATO composite fiber tended to decrease. In particular, the PU–ATO (1) composite fiber showed an IR transmission of ~20%, whereas a very low IR transmission of ≤3% was found for the PU–ATO (2) composite fiber and other specimens. Therefore, for the subsequent textile fabrication and IR- and thermal radiation-shielding experiments, we selected the PU–ATO (3) composite fiber because of its optimum mechanical properties and IR transmission.

As shown in Fig. 3g, h, the PU–ATO (3) composite fiber-based textile showed a hydrophobic property with water contact angles ≥130° in all areas; accordingly, water droplets on the surface of the textile were not absorbed but easily rolled off. On the other hand, the PU–ATO (3) textile not subjected to hydrophobic treatment absorbed water droplets, which remained on the textile until natural drying. The PU–ATO composite fiber is hydrophobic due to the perfluoroalkyl chains of HDF-PA SAMs attached to its surface; the hydrophobicity of the fiber is essential for the fiber to maintain a certain level of IR- and thermal radiation-shielding capability. When the fiber is exposed to water, it absorbs water, and the absorbed water causes IR image distortion according to the absorption of the IR and thermal radiation. Accordingly, the pristine PU–ATO (3) composite fiber-based textile showed IR distortion in all areas where water was absorbed, whereas the hydrophobic PU–ATO (3) composite fiber-based textile could maintain stable IR- and thermal radiation-shielding properties because it was not wetted by water (Supplementary Figs S6, S7, Supplementary Information). Figure 3i shows the IR- and thermal radiation-shielding properties after the removal of water droplets from hydrophobic and nonhydrophobic (untreated) PU–ATO (3) composite fiber-based textiles. After placing pristine and hydrophobic PU–ATO (3) composite fiber-based textiles, each with a size of ~50 mm², side by side on a hot plate heated to 40 °C, we observed the textiles with a thermal IR camera after 1 min. The results indicated that the hydrophobic PU–ATO (3) composite fiber-based textile exhibits stable IR- and thermal radiation-shielding properties, whereas the PU–ATO (3) composite fiber-based textile not subjected to hydrophobic treatment exhibits different IR and thermal radiation intensities depending on whether the textile is dry or wet. This confirms that IR and thermal radiation intensity could be distorted by water.

Figure 4 compares the IR- and thermal radiation-shielding properties of the PU–ATO (3) textile when irradiated with thermal IR light in the high- and low-temperature range. Figure 4a shows the optical and IR images of polyester, PU, and PU–ATO (3) composite
Textiles placed on top of the hot plate, a thermal IR light source. Changes in the temperature of three types of textiles (~50 mm²) placed 2 mm above the surface of the hot plate heated to 40 °C were observed over a period of 120 min with a thermal IR camera. Note that the temperature at a distance 2 mm away from the 40 °C hot plate was ~36 °C. For each textile, the temperature from nine different points on the textile was measured, and the mean value with standard deviation is reported. The textiles of different materials placed on top of the hot plate showed different patterns of surface temperature over time. As the polyester textile has weak IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min. Further, because of its inherent IR- and thermal radiation-shielding properties, its surface temperature reached ~35.0 °C immediately (0 min) after it was placed 2 mm from the surface of the hot plate at 40 °C, and the same temperature appeared even after 120 min.
condition of −20 °C, the change in the temperature of the PU–ATO (3) composite textile was modest in comparison with those of the polyester and PU textiles (Supplementary Fig. S9, Supplementary Information). Thus, the results indicated that the IR- and thermal radiation-shielding properties of the PU–ATO (3) textile function effectively under both high- and low-temperature conditions.

Figure 4d shows the surface temperature reached by the polyester, PU, and PU–ATO (3) composite textiles after 60 min under varying applied temperature conditions, including room temperature (20 °C) as the reference and −20, −3, +30, +40, +50, +60, +70, and +80 °C. The polyester and PU textiles attained a surface temperature that was as high as ~90% of the applied temperature. In contrast, the PU–ATO (3) composite textile showed very small changes in the surface temperature, despite the significant changes in the applied temperatures. Under the harshest temperature condition of 80 °C, the polyester and PU textiles showed surface temperatures of −8.1 and −1.3 °C, respectively, with differences of −11.9 (polyester) and −18.7 °C (PU) compared with the applied temperature (−20 °C). In contrast, the PU–ATO (3) composite textile with strong IR- and thermal radiation-shielding properties showed surface temperatures of 45.1 °C and 9.1 °C, respectively, under 80 °C and −20 °C conditions. The results indicate that despite the applied temperature being changed by 100 °C (−20 to +80 °C), the surface temperature of the PU–ATO (3) composite textile changed by only 36.0 °C, which is only ~1/3 of the actual change in applied temperature. For reference, under the condition of 30–40 °C, which is similar to human body temperature, the surface temperature of the PU–ATO (3) composite textile was 24–27 °C. This temperature is lower than that of polyester (29–37 °C) and PU (28–35 °C), indicating the more effective shielding of IR and thermal radiation by the PU–ATO (3) composite textile.

To be used as a practical wearable textile, the textile must maintain its IR- and thermal radiation-shielding properties in an environment with a constantly changing temperature. For this, the reliability of IR- and thermal radiation-
shielding properties must be verified by repeatedly applying high- and low-temperature conditions or by considering its performance during long-term exposure under a single temperature condition. Figure 5a shows the IR images of polyester-based, PU-based, and PU–ATO (3) composite-based textiles after exposure for 60 min to temperatures of 80 and −20 °C. Note that a hot plate was used to apply the high temperature of 80 °C, and a Peltier module was used to apply the low temperature of −20 °C. In the IR images, the PU–ATO (3) composite textile shows a lower temperature of ~45 °C under the high-temperature condition of 80 °C. In contrast, it shows a temperature of ~9 °C under the low-temperature condition of −20 °C, which is higher than that of the polyester and PU textiles. Figure 5b shows the results of repeatedly exposing the PU–ATO (3) composite textile to high (+80 °C) and low (−20 °C) temperatures. The surface temperatures were measured after 60 min at each of the high- and low-temperature conditions, with the cycle repeated ten times. The surface temperature of the PU–ATO (3) composite textile remained steady at 45.8 ± 0.6 °C and 8.8 ± 0.4 °C after repeated exposure to high (+80 °C) and low (−20 °C) temperatures, respectively. These findings confirm that the IR- and thermal radiation-shielding properties of the PU–ATO (3) composite textile could be maintained even when repeatedly exposed to high- and low-temperature conditions. The radiation absorbed by the surface of the object is converted into thermal energy through the vibration of the molecules, thereby increasing the surface temperature of the object and emitting radiant heat corresponding to the fourth power of the surface temperature through infrared rays. The heat radiation energy, referred to as thermal infrared rays, can thus be controlled according to the temperature and wavelength. As such, the infrared rays have a deep correlation with heat and cannot be explained separately from heat. Therefore, the PU–ATO (3) composite textiles with low heat-transfer capability are able to extensively block IR radiation. Figure 5c shows the continuous IR- and thermal radiation-shielding properties of the PU–ATO (3) composite textile over time. The applied temperature was set to 40 °C in consideration of the typical environment encountered. The surface temperature of the textile was measured with application of a temperature of 40 °C for 8 h per day for a total of 30 days. The surface temperature of the textile was maintained at ~25 °C, even over a long duration. Furthermore, Fig. 5d shows the IR- and thermal radiation-shielding properties of the PU–ATO (3) composite textile over different durations (1, 3, 7, 14, 20, and 30 days). As shown in the IR images, the IR- and thermal radiation-shielding properties were maintained over time.

Conclusion
In this study, we fabricated a PU–ATO composite fiber and used it to weave wearable textiles with IR- and thermal radiation-shielding properties. The ATO sol prepared by a typical sol–gel method showed changes in its IR transmission property according to the process conditions, such as the aging time and concentration. Moreover, a PU–ATO composite fiber consisting of an ATO sol with IR- and thermal radiation-shielding properties eveny embedded in a PU matrix was fabricated, and the mechanical, chemical, and optical properties of the obtained fabric were analyzed. Based on the findings, the optimal concentration and aging time of the ATO sol mixed with PU for a wearable textile were identified. The PU–ATO (3) composite fiber used to fabricate IR- and thermal radiation-shielding textiles had a tensile strength of 3.90 MPa and strain of 335.7%; the ATR–FTIR study confirmed the IR transmission to be ≤2%. The IR- and thermal radiation-shielding textile made from optimized PU–ATO (3) composite fiber showed excellent IR- and thermal radiation-shielding properties when observed with a thermal imaging camera applied to heating and cooling sources (−20 to 80 °C), such as a human body, hot plate, or Peltier cooling plate. Moreover, a hydrophobic treatment was performed on the fabricated PU–ATO composite fiber to prevent the distortion of the IR- and thermal radiation-shielding properties due to the absorption of water. To assess the reliability of the PU–ATO composite fiber, it was exposed repeatedly (ten times) to extreme temperature conditions of 80 °C and −20 °C and subjected to long-term exposure at 40 °C for 30 days. The results confirmed that the fiber’s IR- and thermal radiation-shielding capabilities can be stably maintained under those conditions. With the continuing development of stealth technology used to conceal IR signals originating from objects, the PU–ATO composite fiber is expected to lead to a new market surrounding wearable IR stealth products; such products can advantageously be fabricated into a wearable form, which is not feasible with conventional thin films or paint coating methods.

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