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Investigating polymer infiltration kinetics in nanoporous metal scaffolds using UV-vis spectroscopy†

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This work demonstrates that the optical response of nanoporous gold (NPG) is directly related to the infiltration extent (IE) of the polymer. Infiltration of poly(2-vinylpyridine) (P2VP) into NPG with a pore radius (R_p) of 34 nm was investigated at 140 °C. The UV-vis spectra of the NPG display plasmon absorption peaks due to the Au ligaments that depend on the height of the P2VP infiltration front in the NPG film. During infiltration, the absorption peak position shifts to longer wavelengths, while the peak height, width, and area increase monotonically with time. For P2VP with molecular weights (M_w) from 85 kDa to 940 kDa, the time to reach 80% of the IE ($\tau_{80\%}$) scales as $M_w^{1.35}$, in good agreement with *in situ* spectroscopic ellipsometer results. AFM and XPS support the strong attraction of P2VP for the Au surface and pores as demonstrated by wetting of P2VP over surface ligaments and a shift of the 4f orbital from the N on P2VP to higher binding energy, respectively. Using nanorods configured as a “T” to model ligament geometry, discrete dipole approximation (DDA) simulations capture the optical properties of the P2VP/NPG nanocomposite during infiltration and confirm experimental results. The evolution of the P2VP/NPG optical properties is attributed mainly to an increase in the effective refractive index within the pores. This study presents UV-vis spectroscopy as an alternative method for studying polymer infiltration into nanoporous metal scaffold films.

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Introduction

Polymer nanocomposites (PNCs) have garnered significant attention due to their unique combination of organic and inorganic components,^{1–3} leading to a range of applications as coating,^{4,5} membranes^{6,7} and actuators.⁸ PNCs with a high inorganic nanofiller content have received much attention because of their ability to improve mechanical properties.^{9,10} However, creating PNCs with a high loading of nanofiller (>50 vol%) is challenging because the thermodynamics of mixing typically favor nanofiller aggregation and as a result undesirable materials heterogeneity. Various approaches have been developed to enhance nanofiller loading capacity and distribution using pre-fabricated scaffolds, *in situ* polymerization, nanoparticle assemblies. Examples include the infiltration of polymers into anodized aluminum oxide (AAO)

membranes,^{11–16} *in situ* polymerization with graphene-based nanofillers,^{17,18} nanoparticle assemblies,^{19–29} and nanoporous gold (NPG).^{30,31} In our previous studies,^{30,31} we fabricated polymer-infiltrated NPG (PING), with a high nanofiller loading of approximately 50 vol%, by infiltrating a polymer melt into a nanoporous gold scaffold. During the fabrication of the PING composite, polymer kinetics were investigated under conditions of moderate confinement, where chain size was slightly less than the pore diameter. Recent attention^{11,13,15,30–46} has been given to polymer kinetics under confinement because spatial limitations can alter polymer conformations, thus affecting thermal properties and melt viscosity. For instance, studies have shown that polymers inside channels exhibit an enhanced glass transition temperature (T_g) as compared to the bulk.^{23–25,30,47} Infiltration kinetics has garnered special interest due to its potential to precisely control properties such as optical response, mechanical performance, and surface topography through careful management of the extent of polymer infiltration.^{25–27,30,31} Research on polymer kinetics in confined channels is important for fabricating new membranes that respond to stimuli and facilitate ion conductivity.

To quantify polymer infiltration into porous media, several techniques have been used, including cross-section SEM,¹⁶ small

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angle X-ray scattering (SAXS),¹³ dielectric spectroscopy (DS),^{11,12,14,15,48–51} and spectroscopic ellipsometry (SE).^{23–28,30,31,47} Cross-section SEM can directly image polymer height inside confined pores. At 130 °C, Hu and Cao studied the infiltration of a polyethylene (PE) melt inside AAO templates (length = 60 μm, diameter = 200 nm).¹⁶ The kinetics of infiltration was determined by measuring the PE growth front position (H) as a function of infiltration time.¹⁶ The infiltration height was proportional to the square root of infiltration time,¹⁶ such that $H(t) \sim 0.28t^{0.5}$, where the units of H and t are 10^{-6} m and s, respectively. This scaling is consistent with the Lucas–Washburn equation, indicating that the PE melt within the AAO behaves like a simple fluid.¹⁶ Drawbacks of SEM include the need for image analysis to locate the growth front, and polymer degradation under the electron beam.⁵²

SAXS has also been used to monitor polymer infiltration.¹³ Shin *et al.* utilized time-resolved SAXS to observe the capillary rise of polystyrene (PS) within an AAO membrane.¹³ In this approach, the integrated intensity (Q) was used to extract the volume fractions of pore (Φ_p) and alumina (Φ_{AO}) using $Q \sim [\Phi_U(\rho_{AO})^2 + \Phi_F(\rho_{AO} - \rho_{PS})^2]\Phi_p\Phi_{AO}$, where ρ is the electron density, and Φ_F and Φ_U are the volume fraction of filled and unfilled pores such that $\Phi_F + \Phi_U = 1$.¹³ The total flux (J) is given by

$$J = \pi r^2 \frac{\Delta l}{\Delta t} = -K \frac{\Delta \left(\frac{Q}{Q_0} \right)}{\Delta t} \quad (1)$$

where r is the radius of the pore and l is the length that the polymer has traveled into the membrane. In eqn (1), the flux is related to the change in Q through $K = \pi r^2 L(\rho_{AO})^2 / (2\rho_{AO} - \rho_{PS})\rho_{PS}$, where L is membrane length. As pores fill with polymer, the integrated scattering intensity decreases from Q_0 to Q .¹³ Using the preceding equations, Φ_U can be calculated. While SAXS was one of the first methods to measure infiltration, it has several limitations, including access to X-rays, polymer degradation,⁵³ and low intensity due to membrane scattering.

Dielectric spectroscopy (DS) is used to measure the dielectric properties of materials. A layer of gold acts as the upper electrode, while the substrate serves as the lower electrode.^{54–56} DS measures the complex dielectric permittivity (ϵ^*) as a function of frequency (ω) and temperature (T). The measured $\epsilon^*(\omega, T)$ are fitted through the Havriliak and Negami (HN) equation:

$$\epsilon_{HN}^*(\omega, T) = \epsilon_\infty(T) + \sum_{k=1}^2 \frac{\Delta\epsilon_k(T)}{[1 + (i\omega\tau_{HN,k}(T))^{m_k}]^{n_k}} + \frac{\sigma_0(T)}{ie_f\omega} \quad (2)$$

where $\Delta\epsilon_k(T)$ is the relaxation strength, $\tau_{HN,k}(T)$ is the relaxation time of asymmetric shape parameter m_k and spectra broadness n_k ($m_k n_k \leq 1$, $m_k > 0$), and ϵ_∞ is the dielectric permittivity at the high-frequency limit. The infiltration time¹⁴ is determined from the relaxation time at maximum loss (τ_{max}).

$$\tau_{max,k} = \tau_{HN,k}(T) \sin^{-\frac{1}{m}} \left(\frac{\pi m_k}{2(1+n_k)} \right) \sin^{-\frac{1}{m}} \left(\frac{\pi m_k n_k}{2(1+n_k)} \right) \quad (3)$$

Previous studies^{11,12,14,15,48–51} showed that DS is a reliable method for tracking polymer kinetics inside a confined channel. However, the application of DS to study infiltration presents some challenges. For example, attractive interactions between

the infiltrating polymer and the pore surface can introduce interfacial effects, which broaden the relaxation processes, making it difficult to decouple polymer dynamics from interfacial dipole contributions. Additionally, in the case of NPG, the NPG structure has a distribution of pore sizes and local environments, leading to a range of polymer relaxation times, which causes uncertainty in τ_{max} .

Spectroscopic ellipsometry (SE) has been widely utilized for measuring polymer infiltration.^{25,26,30,31} SE detects the change in light polarization in the s-plane and p-plane after light reflects from the front and back surfaces of a film.⁵⁷ The polarization change is represented by the amplitude ratio Ψ and phase difference Δ . Optical properties, such as refractive index, extinction coefficient, and optical anisotropy, can be obtained using models to fit the Ψ and Δ at various wavelengths.⁵⁷ SE has been used in our previous infiltration studies of polystyrene (PS) and poly(2-vinylpyridine) (P2VP) into NPG.^{30,31} Because a laser is the probing radiation, the sample heating chamber does not require a vacuum like in SEM and SAXS. Thus, SE lends itself to *in situ* measurements. Disadvantages include modeling the infiltration front as a sharp boundary and the use of fitting parameters to determine Ψ and Δ , which can introduce uncertainties that depend on the model and parameter constraints.

In this manuscript, we present UV-vis as a reliable and facile tool to measure polymer infiltration into nanoporous gold (NPG). Specifically, the localized surface plasmon resonance (LSPR) can be used to characterize PNCs containing metallic nanoparticles (NPs) as well as metallic nanoporous membranes. For Au NPs and membranes, the LSPR adsorption frequency depends on the separation of AuNPs (ligaments), as well as their orientation⁵⁸ and volume fraction.⁵⁹ For metallic films, the LSPR depends on film thickness and valence electron density.⁶⁰ For spherical Au NPs, the LSPR is in the visible range,⁶¹ and undergoes a slight red shift in peak position as particle diameter increases. For Au nanorods, the LSPR exhibits longitudinal and transverse modes due to oscillations of free electrons along each axis.⁶² Whereas the transverse LSPR is relatively insensitive to the aspect ratio (AR) of nanorods, the longitudinal LSPR increases as the aspect ratio increases.^{63,64} In contrast to discrete Au NPs, the NPG membrane exhibits a bicontinuous structure of gold ligaments.^{31,65} The NPG exhibits unique optical properties^{66–70} because the bicontinuous structure limits the motion of the conducting electrons⁷¹ resulting in LSPR absorption in the visible range.⁷² The optical response depends on ligament aspect ratio⁷⁰ and pore diameter.⁷² Detsi *et al.* showed that the transmittance peak wavelength position increases from 560 to 580 nm as AR of the ligaments increases from 1.2 to 1.8.⁷⁰ In this study, Detsi *et al.* modeled the optical response of the Au ligaments using a single Au nanorod with an AR from 2.5 to 9.5. Notably, the two plasmonic modes in NPG arise from longitudinal and transverse LSPR in the ligaments rather than resonant absorption from the film thickness. Lang *et al.* observed a red shift in the LSPR as pore diameter increases.⁷² Previous studies on unfilled NPG and water-filled NPG provide important background for our investigation of the



optical response of NPG when the pores are partially and completely filled with polymer.

There are considerations before applying UV-vis to study polymer infiltration into nanoporous scaffolds. First, the nanoporous scaffold must be metallic and exhibit a localized surface plasmon resonance (LSPR). Second, typical polymers do not absorb light in the same range as the LSPR. As a result, the absorbance of polymers is generally separated from that of the metallic ligaments in the scaffold, which typically absorb light in the 500 to 1000 nm range, depending on the size and aspect ratio of the ligaments.⁷⁰ One caveat is that polymers with conjugated backbones, such as polythiophene,⁷³ absorb light in visible or near-IR range. For polymers that have absorbance overlapping with the LSPR, spectrum deconvolution can be performed when applying the UV-vis method. The UV-vis method has one significant advantage compared to spectroscopic ellipsometry (SE). Namely, SE is limited to study ~ 100 nm thick metallic scaffolds because SE relies on measuring the change in polarization by light reflected from the nanocomposite films.^{30,31} In contrast, because UV-vis measures the transmission and absorbance of light, micron-thick NPG films can be characterized. In general, UV-vis spectroscopy can serve as a complementary technique to the more commonly used SE measurements.

Here, we measure the optical properties of NPG during polymer infiltration using UV-vis spectroscopy. The initial sample geometry is a bilayer of a NPG foil (*ca.* 120 nm) over a poly(2-vinyl pyridine) (P2VP) film (*ca.* 120 nm) deposited on a glass substrate.³⁰ During infiltration, the absorbance peak redshifts from 517 nm to 521 nm, increases in intensity and broadens. The infiltration extent (IE) of P2VP into the NPG is determined from the peak height, width, and area. In our previous study of P2VP,³¹ the scaling of IE with molecular weight M_w was measured using *in situ* SE. Using UV-vis spectroscopy, the time for 80% IE is found to scale as $M_w^{1.35}$, in good agreement with SE results.³¹ Using AFM and XPS, the strong attraction of P2VP for Au is evidenced by the observed wetting of P2VP on the NPG surface and a shift to higher binding energy of the 4f orbital of N in the 2VP monomer. Using nanorods arranged in a “T” formation to model ligament absorption, discrete dipole approximation (DDA) simulations more accurately capture the optical properties during infiltration than a single rod. Changes in the optical properties of P2VP/NPG during infiltration are primarily due to an increase in the effective refractive index within the pore. This study presents UV-vis spectroscopy as a facile method to quantify polymer infiltration into a nanoporous metal scaffold. Understanding the optical responses of nanoporous metallic scaffolds during and after polymer infiltration allows for the precise design of partially and completely filled polymer composites, which can potentially be applied as membranes for separations.

Materials and methods

Preparation of nanoporous Au/P2VP bilayer

Poly(2-vinyl pyridine) (P2VP) is dissolved in butanol ($\geq 99.9\%$, Sigma-Aldrich), and stirred overnight. P2VP polymers with

narrow polydispersity are used as received. The radius of gyration (R_g) is calculated by $R_g = \sqrt{\frac{b^2 N}{6}}$, where the Kuhn length (b) is 1.8 nm, the number of Kuhn segments (N) is $\frac{M_w}{M_0}$, and the Kuhn monomer molar mass is 720 Da.⁷⁴ The confinement parameter is defined as the ratio of R_g to R_{pore} , where $R_{\text{pore}} = 34$ nm as characterized in the previous study.⁷⁵ The molecular weight, radius of gyration and confinement parameter for P2VP are shown in Table 1.

The P2VP solutions (2.4 wt%) are spin-coated (4000 rpm, 60 seconds, Laurell Technologies) onto 1×1 cm² cleaned glass substrates and dried at 70 °C for 10 minutes to remove residual solvent. All films were smooth and homogeneous as-cast. Film thickness was measured with a white-light reflectometer (Filmetrics F3-UV) and ranged from 80–150 nm. P2VP is infiltrated into nanoporous gold (NPG) with pore radius (R_{pore}) of 34 nm and thickness of 120 nm, as characterized in our previous publication.⁷⁵ The NPG membrane is deposited on top of the P2VP film. The resulting bilayer is then dried at 70 °C on a hot plate for 5 minutes to remove residual H₂O. For more information about the preparation of the bilayer samples, please refer to our earlier publications.^{30,31} These bilayers are annealed on a Mettler heating stage at 140 °C under consistent Argon flow and then characterized by UV-Vis spectroscopy. The precision of the Mettler heating stage is 0.01 °C.

Ultraviolet-visible (UV-vis) spectroscopy

Absorbance spectra are obtained using a Varian Cary Win 5000 UV-VIS-NIR Spectrophotometer with a wavelength range of 200–1000 nm. Measurements of the absorbance are taken with a frequency of 2 nm per sweep at each infiltration time. The P2VP:NPG bilayer, atop a glass substrate, is annealed at 140 °C for different annealing times until complete infiltration is observed. The absorbance spectra of glass are included in the ESI,[†] and are subtracted from the bilayer absorbance spectra during data analysis.

Atomic force microscopy (AFM)

Surface topography of the P2VP:NPG before and after full infiltration are measured using AFM. Tapping mode AFM is performed using Bruker Icon AFM with tips (TAP300AL-G-50 radius of curvature < 10 nm, Ted Pella). The images collected are 1×1 μm^2 . The surface properties images are processed using Gwyddion software.

In situ spectroscopic ellipsometry (SE)

Spectroscopic ellipsometry (SE) (J.A. Woollam, Alpha SE) is used to measure the refractive index of the P2VP bilayer/

Table 1 Characteristics of the P2VP infiltrated into the NPG

Sample	M_w (Da)	R_g (nm)	$\Gamma = R_g/R_{\text{pore}}$
P2VP-85k	85 000	7.98	0.23
P2VP-302k	302 000	15.05	0.44
P2VP-643k	643 000	21.96	0.65
P2VP-940k	940 000	26.55	0.78



composite during the infiltration process. The SE wavelength range is 380–900 nm. A Linkam THMSEL350V heating stage is used for heating. The precision of the heating stage is 0.1 °C with respect to the set temperature. The heating rate is 30 °C min⁻¹. The samples are heated to 70 °C and held at this temperature for 5 minutes to ensure system equilibrium. Then, the sample is heated at a rate of 30 °C min⁻¹ and then held at 140 °C to study infiltration. Because the bulk P2VP has T_g around 100 °C, temperatures higher than 100 °C will induce polymer infiltration into NPG. As a result, the infiltration time refers to the time after the heating plate reaches 100 °C. The heating stage takes *ca.* 1.3 min to ramp from 100 °C to 140 °C. During infiltration, the temperature is stably held at 140 °C. Effective medium approximation (EMA) Model with two material constitutes is used to capture the change in refractive index within the NPG as polymer fills the pores and approaches the top surface. Details of SE modeling are presented in our previous publications.^{30,31} The infiltration extent (IE) is given by $IE = (n_t - n_i)/(n_f - n_i)$. The initial refractive index, n_i , the refractive index prior to the P2VP infiltration and the final refractive index, n_f , is the refractive index after complete infiltration.

X-ray photoelectron spectroscopy (XPS)

XPS is used to quantify the binding energy of elements in NPG and NPG with polymer before and after it is completely infiltrated. X-ray photoelectron spectroscopy (XPS) was performed in an ultrahigh vacuum chamber equipped with a hemispherical electron energy analyzer (Leybold–Heraeus) collecting at all angles. The incident energy of the Al-K α X-ray source is 1486.6 eV (VG Microtech). The sampling depth is about 3 nm. To provide insight into the attraction at the P2VP/Au interface, XPS is used to quantify the shift in binding energy of the 4f electrons of Au.

Discrete dipole approximation (DDA) simulations

The discrete dipole approximation (DDA) is used to determine the optical properties of the unfilled, partially filled and filled NPG. The MATLAB toolbox containing DDA with surface interaction (DDA-SI) is used.⁷¹ The optical properties are the extinction coefficient, localized surface-plasmon resonance (LSPR) wavelength, and LSPR absorption strength, calculated using the MATLAB toolbox.^{71,76} Because P2VP and air do not exhibit plasmonic resonances and have negligible absorption in the visible range, the dominant contribution to the optical response from the composite is the LSPR from the Au ligaments. Although the dielectric properties of the polymer within the pores influence the plasmonic response, their contribution to direct absorption is minimal over the wavelength range of interest. Thus, in our simulations, we model only the Au ligament structure to represent the NPG. A T-shape structure formed by two nanorods more accurately captures the experimental LSPR absorption spectra compared to a single nanorod, as illustrated in Fig. S13 (ESI[†]). Fig. 1 shows the T-shape structure used to model the NPG and the incident UV-vis radiation propagating in the z direction. This structure captures the plasmon hybridization of the unfilled NPG as well as the

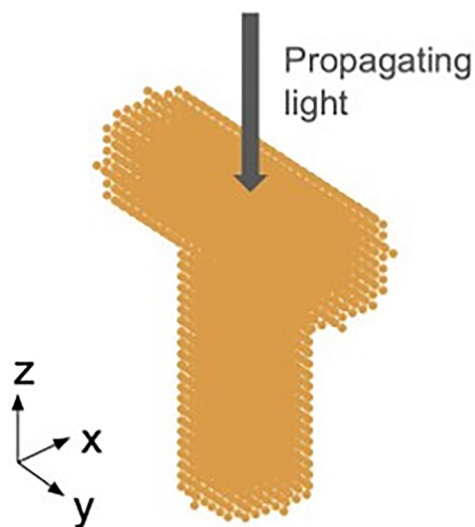


Fig. 1 The T-shaped nanorods represent the ligaments in the NPG and are used in DDA simulations. The individual points represent dipoles excited by the incident radiation. Each dipole is assumed to be dielectric but non-magnetic. All dipoles have identical electric permittivity. For each nanorod, the radius and length are 25 nm and 125 nm, respectively. There are 142 890 dipoles in the T-shaped structure.

reduction in intensity upon infiltrating a polymer into the pore.^{77,78} As shown in Fig. 1, the nanorods have a radius of 25 nm, a length of 125 nm, and aspect ratio of 2.5. These dimensions match the size of the ligaments in the experimental system.

Results and discussion

Transmittance spectra of NPG filled with air, water and P2VP

Plasmon absorption in the ligaments of the NPG depends on dielectric constant within the pore. The transmittance of unfilled NPG, water-filled NPG and P2VP-filled NPG was compared at constant pore radius, 34 nm. The NPG:P2VP nanocomposite was fabricated by annealing a NPG/P2VP bilayer at 140 °C for 3 h to achieve complete filling.³⁰ Fig. 2 compares the room temperature transmittance spectra for NPG:air, NPG:water and NPG:P2VP. The transmittance minimum for NPG:air is weak because of the high reflectivity of the gold in the visible region, which reduces the absorbed (A) and transmitted (T) intensities. When the absorbance is weak (Fig. S9, ESI[†]), the transmittance spectra better represent the optical response because $T = 10^{-A}$.

At the transmission minimum, the intensity decreases and the position redshifts as the refractive index of the medium in the pore increases. Fig. 2 shows that the NPG transmittance spectra minima (*i.e.*, absorbance peak maxima) exhibits a redshift. For the unfilled pores, the minimum is weak and appears at 514 nm. For NPG:water, the transmittance peaks become stronger relative to the minimum. Compared to air ($n = 1$), the NPG has a lower reflectivity when filled with water ($n = 1.33$ @ wavelength = 528 nm).⁸⁰ Because the refractive index difference



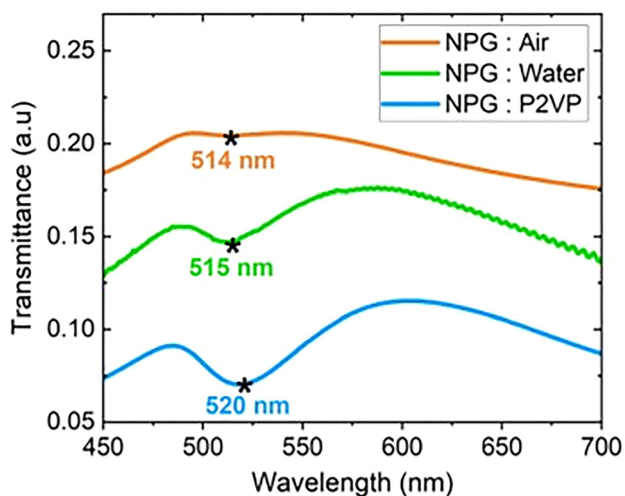


Fig. 2 Transmittance of NPG filled with air (unfilled), water and P2VP at 25 °C. The asterisk denotes the shift of the transmittance peak minimum from 514 to 515 to 520 nm as the pore media changes from air (orange) to water (green) to P2VP (blue). The oscillations in NPG:water result from the background subtraction of the Kapton tape.⁷⁹ Data are shifted to facilitate comparison of spectra.

between gold and water is smaller than gold and air. The NPG:water also undergoes a small red shift from 514 nm to 515 nm due to the increase in the refractive index in the pores. The oscillations at higher wavelengths are due to the background subtraction from the Kapton tape (Fig. S9, ESI[†]).⁷⁹ For NPG:P2VP ($n = 1.54$), the transmittance peaks are stronger than those observed in the NPG:air and NPG:water systems because the reflectivity from the NPG decreases further. For NPG:P2VP, the transmittance minimum appears at 520 nm, indicating a red shift compared to air and water. Interpreting the transmission spectra in terms of absorption, these studies show that the absorption peak exhibits a red shift and grows in intensity as the refractive index of the media in the pores increases.

Absorbance spectra during P2VP infiltration into NPG

As P2VP infiltrates the NPG film from below, the refractive index of the NPG composite increases with time. Motivated by the previous section, which explored how the transmission spectra of nanoporous gold (NPG) are influenced by the medium filling the pores, we investigated the absorbance spectra of NPG during the infiltration of P2VP-643k. This was done at a temperature of 140 °C, with annealing times ranging from 0 to 300 minutes, as illustrated in Fig. 3. At 0 min, corresponding to an NPG/P2VP bilayer on glass, the absorbance peak lies at 517 nm, slightly redshifted compared to the NPG:air sample (514 nm, Fig. 2). This shift is attributed to the interface between NPG and P2VP, as discussed in the mechanism section. With increasing time, the peak shifts to 520 nm after 10 min and reaches 521 nm at 300 min (full infiltration). The absorbance peak also broadens and increases in intensity, consistent with an increase in the dielectric constant inside the pores. Similar redshifts in the absorbance peak and intensity increases were observed for NPG:P2VP-85k, NPG:P2VP-302k, and NPG:P2VP-

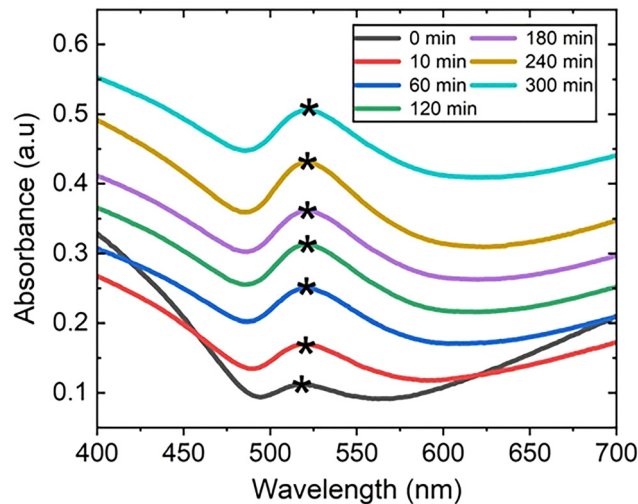


Fig. 3 Absorbance spectra of NPG:P2VP-643k at 140 °C for annealing times from 0 (initial bilayer) to 300 min. The asterisks denote the location of the absorbance peak maximum. Spectra are shifted vertically to better visualize each spectrum.

940k (ESI,† Fig. S10), indicating a common trend during polymer infiltration.

In addition to increasing the dielectric constant in the pore, the absorption peak position and intensity can be affected by strong coupling between the media and the gold surface. For example, the localized surface plasmon resonance (LSPR) of Au shifts to longer wavelengths with stronger intensity when conjugated with poly(vinyl pyrrolidone) (PVP) through electron doping in the valence band.⁸¹ P2VP is expected to be a stronger electron-donor compared to PVP because PVP:Au has a lower Au–Au coordination number than P2VP:Au, namely, 7.5 ± 0.7 and 8.3 ± 0.7 , respectively.⁸² As a result, when P2VP binds to gold, an increase in electron doping in the gold leads to an enhancement of the LSPR, consistent with UV-vis observations. In a later section, the interfacial electron doping of NPG:P2VP will be studied using XPS.

Analysis of absorbance spectra and comparison of infiltration kinetics with SE results

In this section, we relate the absorbance characteristics, namely peak height, peak width and peak area to the extent of P2VP infiltration into the NPG. Using data from Fig. 3, Fig. 4a shows that the relative peak height initially increases rapidly at early times and then more slowly after 100 min. This behavior mimics the increase in pore refractive index measured by *in situ* SE in our previous publication.^{30,31} From the absorption spectra, the initial peak height (h_i) and final peak height (h_f) can be used to determine the infiltration extent, $IE = (h_f - h_i) / (h_f + h_i)$. The time dependence of IE calculated from UV-vis spectra is shown in Fig. 4b. Similarly, the IE was determined from the peak width and peak area as shown in ESI,† Fig. S12. The IE determined from these three approaches (height, width, area) agrees with each other within experimental uncertainty. In the following analysis, only the peak height will be



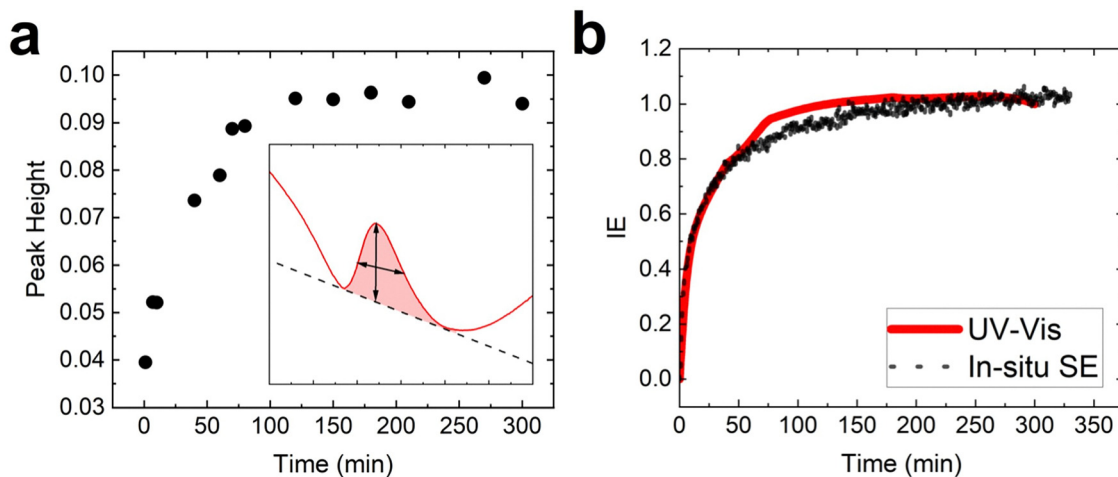


Fig. 4 (a) UV-Vis spectra of absorption peak height versus annealing time for NPG:P2VP-643k at 140 °C. The inset shows the peak height (vertical black arrow), peak width (horizontal black arrow) and area under the peak (highlighted region) from the absorbance curve. (b) Comparison of infiltration extent (IE) for P2VP-643k in NPG measured by UV-vis and SE.

considered. For NPG:P2VP-643k, Fig. 4b compares IE determined by UV-vis (red) and *in situ* SE⁷⁵ (black) methods. The IE values are in good agreement from 0 to *ca.* 0.85. For IE greater than *ca.* 0.85, the UV-vis results exhibit a slightly greater value of IE compared to the SE results. This difference may be attributed to the sensitivity of the plasmon absorption peak to the P2VP wetting the pores ahead of the growth front. Because SE uses a two-layer EMA model, the IE may be slightly less than that measured by UV-vis because it ignores the P2VP wetting ahead of the growth front, which is particularly important near the top surface (*i.e.*, IE > 0.85). At long times (*i.e.*, 300 min), the IE values converge as expected for completely filled NPG. This behavior is consistent with the closing of the gap

between the wetting front (P2VP adsorption on pores) and growth front (P2VP backfilling through coated pores) during late stages of infiltration.

Using data like Fig. 4b for $M_w = 85k, 302k, 643k$ and $940k$ Da, the time to reach 80% IE ($\tau_{80\%}$) is determined and plotted in Fig. 5 for the UV-vis (red) and *in situ* SE (black) methods. The $\tau_{80\%}$ values at each M_w are in statistical agreement according to their standard deviations. In previous studies using *in situ* SE, $\tau_{80\%}$ was found to scale with M_w as 1.40 ± 0.03 , whereas the UV-vis approach yields a scaling of 1.35 ± 0.04 . This good agreement indicates that the UV-vis approach is an accurate method to determine infiltration kinetics in metallic scaffolds. In this section, we demonstrated that UV-vis is a simple method to determine the kinetics of polymer infiltration into NPG and that the results agree with the more common *in situ* SE method.

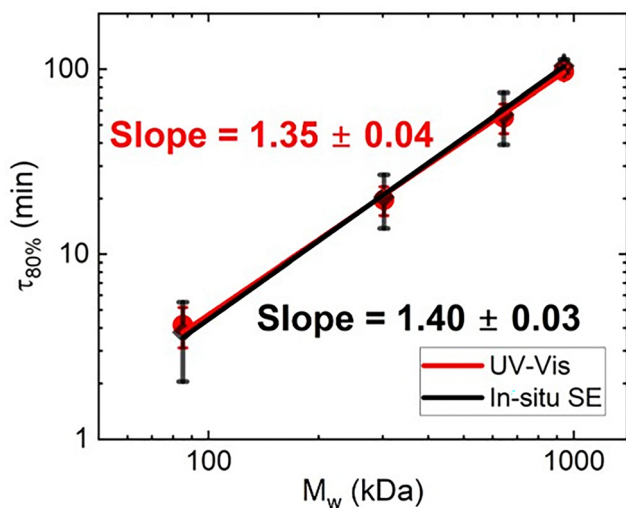


Fig. 5 The 80% infiltration times for P2VP with molecular weights of 85k, 302k, 643k and 940k at 140 °C. The *in situ* SE infiltration times are from our previous study.⁷⁵ The UV-vis (red) and *in situ* SE (black) measurements are in good agreement. Reprinted (adapted) with permission from *Macromolecules*, 2025, **58**(10), 5058–5070. Copyright 2025 American Chemical Society.

Surface characterization of NPG after complete filling by P2VP

The previous section demonstrated that the plasmon resonance peak can be used to quantify polymer infiltration kinetics in a nanoporous metal scaffold. Here, we characterize the near surface region of the NPG:P2VP. For strongly attractive polymer/wall interactions, such as P2VP with gold, simulations show that P2VP forms a strongly adsorbed wetting layer on gold surfaces,⁷⁵ which reduces chain mobility and slows infiltration dynamics. The present study provides new experimental insights into the attraction of P2VP for the Au ligaments.

P2VP forms a thin wetting layer over the top NPG surface after complete infiltration. Fig. 6a and b show AFM height images of the NPG:P2VP-643k surface before and after complete infiltration, respectively. Fig. 6a shows the ligaments (bright) and pores (dark) of the NPG. The length and width of the ligaments are approximately $130 \text{ nm} \times 50 \text{ nm}$, respectively. The RMS roughness is 10.2 nm for the unfilled case. Upon annealing for 5 h at 140 °C to ensure complete infiltration (*cf.* Fig. 3b), the topmost surface of the NPG no longer shows distinct ligaments. This observation differs from polystyrene



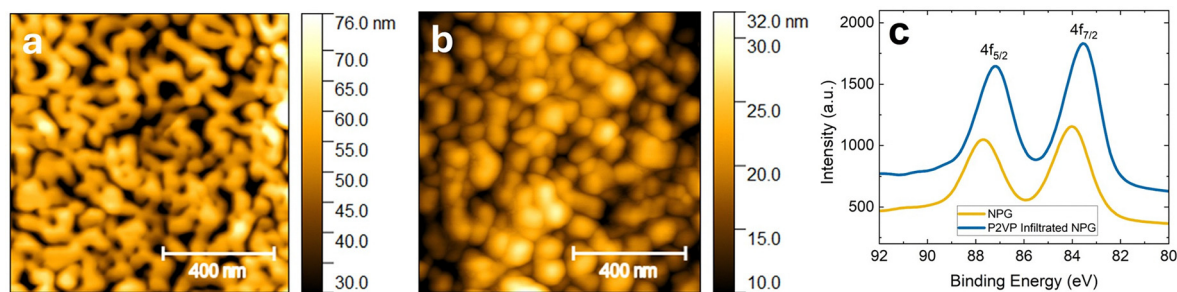


Fig. 6 AFM height images of NPG:P2VP-643k corresponding to (a) an unfilled bilayer ($t = 0$) and (b) a completely filled NPG after annealing at $140\text{ }^{\circ}\text{C}$ for 5 h. Root-mean-square (RMS) roughness values are 10.2 nm and 3.4 nm, respectively. The scanned areas are $1 \times 1\ \mu\text{m}^2$. (c) XPS spectra of Au 4f orbitals for pristine NPG (gold) and the P2VP infiltrated NPG (blue).

(PS) infiltration into NPG where ligaments remain distinct, indicating that PS doesn't completely wet the top surface.³¹ Fig. 6b shows a topography map suggesting that the NPG ligaments are partially covered by P2VP, evidenced by the circular, high features on the surface. Correspondingly, the RMS roughness decreases from 10.2 nm to 3.4 nm, indicating a smoother surface compared to the pre-infiltrated NPG surface. Although the fully infiltrated NPG is less rough than the unfilled scaffold, the surface retains some roughness because of the P2VP droplets that cover the ligaments at the surface (Fig. 6b). The morphology of the completed infiltrated P2VP scaffold agrees with our previous results.⁷⁵ Fig. S1–S3 (ESI[†]) show the corresponding AFM images of P2VP bilayers at $M_w = 85\text{k}$, 302k , and 940k Da before and after complete infiltration. Fig. S4–S7 (ESI[†]) show the corresponding phase images. All images are consistent with a smoother surface after complete infiltration. To confirm that the surface is wet by P2VP, Fig. S8 (ESI[†]) compares the water contact angles (WCA) of fully infiltrated NPG:P2VP (63°) and NPG:PS (70°). These values are consistent with surfaces of pure P2VP and a mixture of PS and Au, respectively, as noted in ESI[†] Fig. S8.

XPS provides insight into the electronic structure of gold atoms in contact with the vinyl pyridine units from P2VP. XPS measures the binding energy of electrons, with a sampling depth of about 3 nm. Fig. 6c shows the binding energies of the core-shell $4f_{5/2}$ and $4f_{7/2}$ electrons before (gold) and after (blue) complete P2VP infiltration into NPG. The pristine NPG has $4f_{5/2}$ binding energy at 87.63 eV, while the $4f_{7/2}$ binding energy lies at 84.02 eV. These values are in good agreement with literature.⁸² Upon infiltration with P2VP, the corresponding binding energies decrease to 87.15 eV and 83.56 eV, a shift of 0.48 eV and 0.46 eV relative to the pristine case. The reduction in binding energies is consistent with a change in the chemical environment spatial redistribution of the valence electrons which create a change in the binding energies of the core electrons. We attribute this shift to electron repulsion in the valence shell, consistent with electron donation from P2VP to Au.^{83,84} The decrease in core-shell binding energies can be attributed to the strong interaction between P2VP and Au. The electron-donating behavior of P2VP originates from the alkyl chains adjacent to the nitrogen atom,⁸² making nitrogen more electronegative, thus enhancing the interaction between the 2VP monomer and

Au at the interface. In summary, the wetting of P2VP on Au ligaments is consistent with the decrease in the binding energy of Au 4f electrons attributed to the attraction between Au atoms and adsorbed P2VP although the change in the UV-vis spectra during P2VP infiltration is mainly due to an increase in the dielectric constant within the pore, the interfacial interactions between P2VP and Au can also affect the LSPR spectra. Further studies are needed to explore the influence of interfacial interactions and NPG absorbance. In the next section, the discrete dipole approximation (DDA) and the effective medium approximation (EMA) are used to model the optical response of NPG during polymer infiltration.

Analysis of absorption spectra of NPG filled with air, water and P2VP

The discrete dipole approximation (DDA) was used to determine the adsorption spectra of unfilled, partially filled, and filled NPG. As shown in Fig. 1, two perpendicular nanorods forming a T were used to capture the Au ligament structure. Although previous research used a single nanorod,⁷⁰ the present studies suggest that a "T" structure could better capture LSPR. The "T" provides a simple representation of the NPG ligament structure while also introducing plasmon hybridization effects, which reduces the sensitivity of the LSPR to changes in the surrounding dielectric medium.⁷⁷ For comparison, the DDA simulations for a single nanorod are presented in Fig. S13 (ESI[†]). The longitudinal LSPR is highly sensitive to polymer infiltration and will be the primary focus of DDA simulations. To test DDA and the "T" model, Fig. 7 shows the absorption spectra of NPG in air ($n = 1.0$), water ($n = 1.33$), and P2VP ($n = 1.54$). As the refractive index in the pore increases from $n = 1.0$ to $n = 1.54$, the peak shifts towards higher wavelengths, namely 515, 521, and 522 nm, respectively. Because the absorbance spectra scale with the inverse of the transmission spectra, we can compare the DDA with experimental results. The DDA spectra are in qualitative agreement with the transmittance spectra from NPG in the same medium as shown in Fig. 2, with peak positions at 514 (air), 515 (water), and 520 (P2VP) nm, respectively. The difference between the UV-vis and DDA spectra near 700 nm is attributed to the scattering contribution in the former. The peak height also increases as the refractive index within the pore increases.



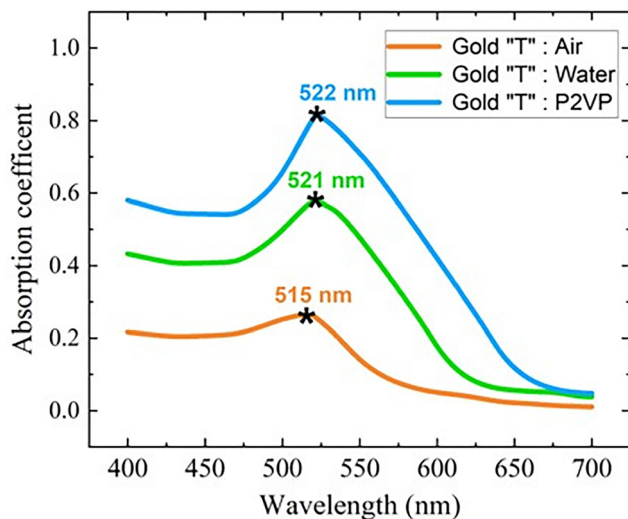


Fig. 7 DDA simulations of the absorption coefficient of NPG for pores filled with air, water and P2VP. As the refractive index within the pore increases the plasmon absorption peak shifts from 515 to 521 to 522 nm. The peak height also increases.

Namely, the peak height increases from 0.264 to 0.581, and then further to 0.815 (arbitrary units), respectively, in qualitative agreement with experimental results shown in Fig. 2. These

simulations show that the absorption peak position and intensity of NPG depend on the refractive index of the media within the pore.

Converting the DDA absorption spectra to infiltration extent of P2VP

To investigate the relationship between P2VP infiltration and absorbance, the optical properties of NPG:P2VP composites at different infiltration extents (IE) were calculated for wavelengths from 400 to 700 nm. As shown in Fig. 8a, the total height (TH) of the T shape is 150 nm (left). The infiltrated height (IH) is set manually, and the surrounding medium has a refractive index of 1.54. IE is given by $100\% \times (IH/TH)$. For example, Fig. 8a (middle) shows a partial infiltration case for IE = 0.5, where bottom region is P2VP (purple) and the top region is surrounded by air ($n = 1.0$). Fig. 8a (right) shows the case of complete infiltration, IE = 1. As IE increases from 0 to 1, Fig. 8b shows that the absorption peak shifts from 515 to 522 nm, and the peak width and intensity broaden and increase, respectively. The simulation results qualitatively agree with the experimental observations in Fig. 3. Noticeably, the left shoulder of the absorption coefficient spectra remains constant at ~ 470 nm during infiltration, in agreement with results in Fig. 3, while the red-shifted right shoulder broadens the peak, consistent with previous experiments.^{63,64} Interestingly, the

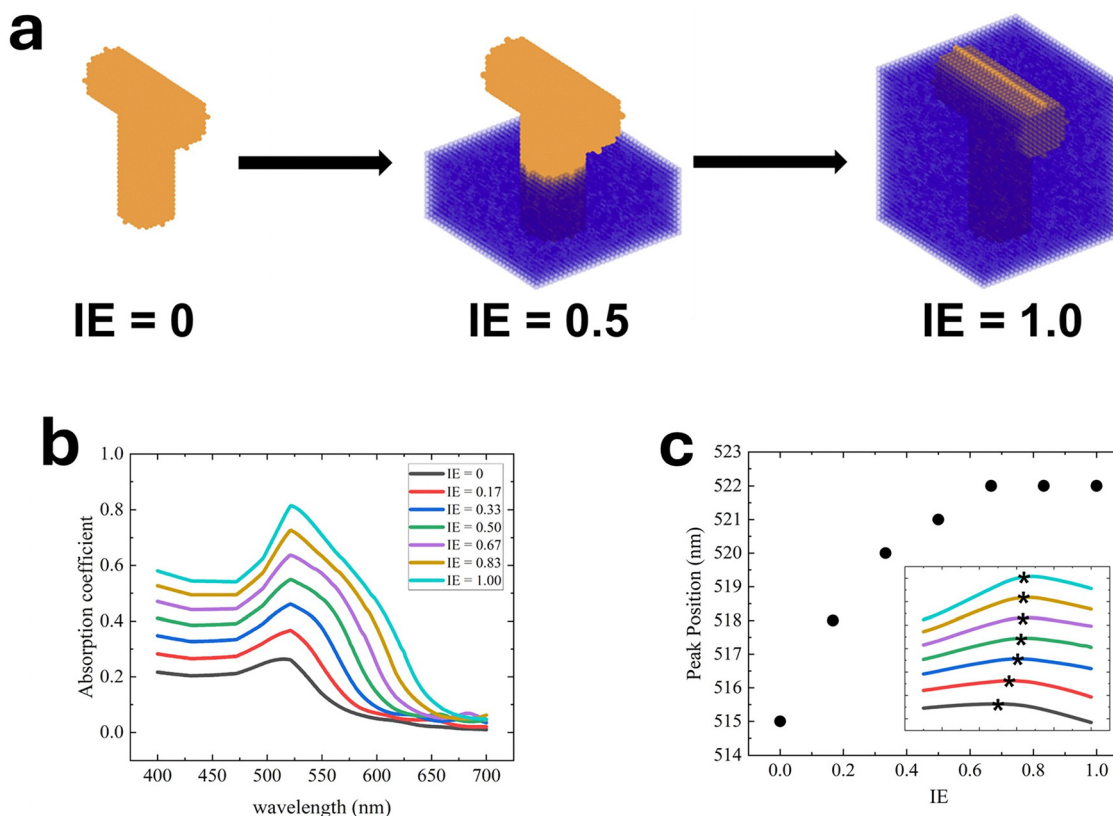


Fig. 8 (a) Schematic of the T-shaped Au nanorod structure (gold) corresponding to infiltration extents (IE) of 0, 0.5, and 1.0. IE is defined as the infiltrated height relative to the total height of the T-shaped structure (150 nm). The medium representing P2VP is shown in purple. (b) DDA calculation of the absorption spectra for unfilled (IE = 0), partially filled (0.17, 0.33, 0.50, 0.67, 0.83), and filled (1.0) states. (c) The peak position versus infiltration extent for unfilled, partially filled and filled states. The inset highlights the peak shift between 490 nm to 540 nm for data in Fig. 8b.



work by Detsi *et al.* also shows a relatively constant left shoulder and a shift in the right shoulder as the NPG aspect ratio (AR) increases.⁷⁰ Namely, as AR increases from 3.8 to 5.1,⁷⁰ the transmittance peak becomes broader and deeper. The left shoulder (transverse) undergoes a blue shift of ~ 3 nm whereas the right shoulder (longitudinal) undergoes a red shift of ~ 32 nm.⁷⁰ The two shoulders in the transmittance spectra originate from the absorption minima in the UV-vis spectra.

The increase in the refractive index of the surrounding medium also accounts for the red shift of the LSPR adsorption peak. Fig. 8c shows the peak position as a function of infiltration extent. Initially, as IE increases from 0 to 0.5, the peak position increase is large, from 515 nm to 521 nm. However, as IE increases from 0.5 to 1.0, the shift is much less, increasing from 521 nm to 522 nm. Interestingly, the peak position approaches a constant value of 522 nm near IE = 0.67, before complete infiltration (IE = 1.0) is reached. However, the adsorption peak width and shape continue to evolve for IE = 0.67, 0.83, and 1.0. The decoupling of peak position and absorption intensity as a function of IE requires further investigation.

Correlation between experimental and DDA simulated infiltration extents

Although the DDA simulations do not explicitly include infiltration time, this section shows that the DDA simulations correlate with UV-vis measurements of IE, which are time-dependent. Whereas the DDA simulation correlates adsorption peak position with IE (Fig. 8c), UV-vis experiments measure peak height as a function of infiltration time, as shown in Fig. 4a. As the height of the medium surrounding the “T” structure increases (*i.e.*, IE increases), the DDA absorption peak position, width, and height increase. As shown in Fig. 3, the UV-vis absorption spectra evolve similarly as a function of time. This observation suggests that the IE from DDA can be correlated with experimental measurements of IE. For the UV-vis method, peak height initially increases rapidly and more slowly as infiltration time increases. As shown in Fig. 4b, the scaling of IE with infiltration time is similar. To build upon the DDA peak position analysis in Fig. 8c, the peak height, full width at half maximum (FWHM), and area under the curve (AUC) are quantified and analyzed. As shown in Fig. 9, the peak height (a.u.) increases linearly from 0.264 to 0.815 as IE increases from 0 to 1 with a slope of 0.55 ± 0.01 . The AUC and FWHM also increase linearly as IE increases (ESI[†], Fig. S15). By using the linear scaling of peak height, AUC, and FWHM as a function of IE, the absorbance spectra can be determined. In summary, DDA simulations of the absorbance spectra show that the peak height, area under the curve (AUC), and full width at half maximum (FWHM) increase linearly with infiltration extent, as illustrated in Fig. 9 and Fig. S15 (ESI[†]). These relationships support the conclusion that changes in the absorbance spectra during infiltration correlate with an increase in the refractive index as the polymer growth front approaches the surface of the “T” structure (DDA) or NPG (UV-vis).

To further investigate the relationship between peak shape and IE, the manually set IE from the DDA simulations (Fig. 8c) are compared with the calculated IE as shown in Fig. 9. Using

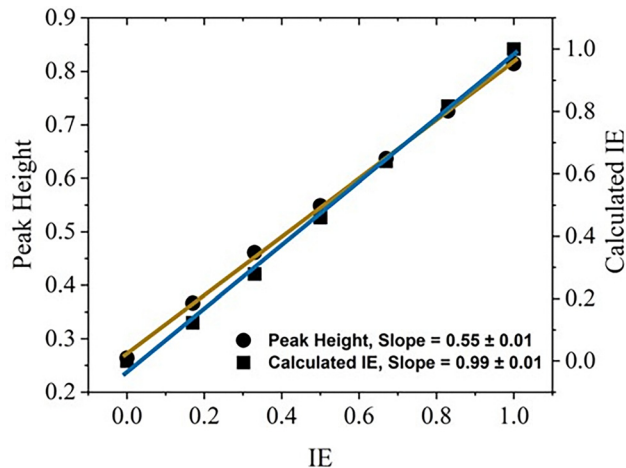


Fig. 9 DDA peak height (solid circles; left y-axis) of “manually” infiltrated T-shaped nanorods (e.g., Fig. 8a) and calculated infiltration extent (solid squares; right y-axis) as a function of infiltration extent (IE). The linear fit for the peak height yields a slope of 0.55 ± 0.01 with Pearson’s $r = 0.99962$ and R -square (COD) = 0.99925. Infiltration extent is defined as the infiltration height of the medium relative to the total height of the “T” model (150 nm). For the DDA spectra, the area under the curve (AUC) and full width at half maximum (FWHM) also increase linearly with infiltration extent, as shown in Fig. S15 (ESI[†]). The correlation between the calculated IE and IE is good, with a slope of 0.99 and a standard error of 0.01.

the peak height (h) values from Fig. 8a, the calculated IE is given by $IE_C = (h - h_0)/(h_f - h_0)$, where $h_0 = 0.264$ and $h_f = 0.815$. The calculated IE increases linearly as IE increases, with a slope of 0.99 ± 0.01 . The agreement between the calculated and manually set IE confirms that the absorbance peak captures polymer infiltration kinetics. Similar comparisons were also conducted for AUC and FWHM, leading to similar results with a linear slope close to 1 (Fig. S16, ESI[†]). This DDA analysis demonstrates that UV-vis absorption spectra can effectively quantify polymer infiltration in a metallic scaffold, like the nanoporous gold used in this study.

Conclusion

This paper demonstrates that UV-vis spectroscopy is a facile method to measure polymer infiltration into nanoporous gold (NPG). At 140 °C, P2VP ($M_w = 85$ –940 kDa) is infiltrated into an NPG film with pore radius of 34 nm. The absorption spectra exhibit a maximum due to the transverse local surface plasmon resonance from the ligaments in the NPG. During annealing, the adsorption peak height and width increase, and the peak position undergoes a red shift. The main reason for this behavior is attributed to an increase in the refractive index within the pores as the P2VP growth front approaches the NPG surface. The peak height, width, and area under the curve (AUC) are found to correlate with the infiltration extent (IE) of P2VP into NPG. UV-vis measurements show that the time for P2VP to reach 80% ($\tau_{80\%}$) infiltration scales as $M_w^{1.35}$, in good agreement with SE results. Moreover, the $\tau_{80\%}$ values at each M_w are in good agreement with SE results. This comparison



shows that plasmonic absorption can be used to measure the kinetics of polymer infiltration in NPG. These studies also show that the absorption characteristics (height, width, AUC) are directly proportional to the IE.

AFM and XPS surface studies support strong wetting of P2VP for the Au surface and pores, as demonstrated by wetting of P2VP over surface ligaments and a shift to higher binding energy of the core shell 4f electrons in Au, respectively. Further studies are needed to quantify the effect of vinyl pyridine adsorption on the surface plasmon resonance of Au. For example, XPS can be used to measure the binding energy of the valence electrons in unfilled NPG and completely filled NPG. Because the valence electrons are the source of the LSPR,⁵⁷ a shift in the binding energy would indicate that the NPG/polymer interface can influence absorption beyond the dominant effect due to the increase in the refractive index (dielectric constant) within the pore.

To model ligaments in the NPG, we use nanorods arranged in a “T” configuration. Discrete dipole approximation (DDA) simulations of absorption are conducted over a range of 400 to 700 nm to capture the transverse localized surface plasmon resonance (LSPR) from the NPG. The infiltration extent is manually set by filling the area around the rod with polymer from the bottom (IE = 0) to the top (IE = 1) of the “T”. The peak shift, height, width and AUC as a function of increasing IE are found to capture UV-vis experiments, even though time is not explicitly included in the DDA simulation. These results show that the effective medium refractive index of the T-shaped nanorods and polymer captures changes in the absorption spectra of the NPG. Whereas this study involves a polymer that strongly absorbs to gold, future studies of hydrophobic polymers will provide insight into the dominant contribution to changes in the absorption peak, namely the refractive index within the pore *versus* the interfacial effects due to polymer adsorption on the pore walls. Grafting polymer onto the pore wall would be an alternative approach to modify the plasmonic properties of the NPG. The present study presents UV-vis spectroscopy as a facile method for studying polymer infiltration into nanoporous metal films. By understanding the optical responses of polymer infiltration of a NPG scaffold, filled and partially filled polymer composites can be precisely fabricated with potential applications as actuators and separation membranes.

Author contributions

R. J. C. and W. K. conceived and planned for the project. C. P. and W. K. developed the experimental procedure and conducted all experimental measurements (besides XPS) and analysis. R. L. and W. K. conducted all simulation development and analysis. M. F. conducted the XPS experiments. W. K., C. P., R. L. wrote the manuscript. R. J. C., and J. M. V. edited the manuscript. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article have been included as part of the ESI.†

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