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Emissive Azobenzenes Delivered on a Silver Coordination Polymer

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ABSTRACT

Azobenzene has become a ubiquitous component of functional molecules and polymeric materials because of the light-induced trans→cis isomerization of the diazene group. In contrast, there are very few applications utilizing azobenzene luminescence since excitation energy typically dissipates via non-radiative pathways. Inspired by our earlier studies with 2,2′-bis[N,N′-(2-pyridyl)methyl]diaminoazobenzene (AzoAMoP) and related compounds, we investigated a series of five aminoazobenzene derivatives and their corresponding silver complexes. Four of the aminoazobenzene ligands, which exhibit no emission under ambient conditions, form silver coordination polymers that are luminescent at room temperature. AzoAEpP (2,2′-bis[N,N′-(4-pyridyl)ethyl]diaminoazobenzene) assembles into a three-dimensional coordination polymer
(AgAAEPpP) that undergoes a reversible loss of emission upon the addition of metal coordinating analytes like pyridine. The switching behavior is consistent with the disassembly-reassembly of the coordination polymer driven by displacement of the aminoazobenzene ligands by coordinating analytes.

INTRODUCTION

Azobenzene (AB) undergoes trans→cis isomerization when irradiated, and cis→trans isomerization thermally or upon exposure to a different wavelength of light. Since light absorption depends on the conjugated π-orbital system, ring substituents change the photoisomerization process and the optical properties of the AB chromophore. The S_2←S_0 transition of aminoazobenzene (aAB) shifts to longer wavelengths, which typically leads to an overlap with the S_1←S_0 transition. As we previously reported, AzoAMoP (1, 2,2'-bis[N,N'-(2-pyridyl)methyl]diaminoazobenzene) exhibits overlapping S_2←S_0 and S_1←S_0 transitions with a λ_{max} at 490 nm and 30-fold higher emission than AB at 77 K. Furthermore, AzoAMoP undergoes minimal trans→cis isomerization due to intramolecular hydrogen bonding between the anilino protons and the pyridyl and diazene nitrogen atoms. Hydrogen bonding imposes an energetic barrier that prevents the aryl rings from adopting the prerequisite collinear conformation necessary for isomerization via the concerted inversion mechanism.

To further develop the photochemistry of these unique aAB derivatives, we replaced the pyridine ligand of the aminomethylpyridine groups with a series of hydrogen bond acceptors, and changed the linker between the amine and pyridine to an ethylene. Investigations with the small library of compounds suggested that reproducing the structure-induced photophysical properties
observed with AzoAMoP would be difficult with a standalone AB chromophore. This study included the report of AzoAEoP (2, 2,2′-bis[N,N′-(2-pyridyl)ethyl]diaminoazobenzene), which utilizes a pyridine ligand and an ethylene spacer. With AzoAMoP and AzoAEoP in hand, we reasoned that using a 3-pyridyl (meta) or 4-pyridyl (para) ligand in the existing aminoazobenzene scaffold would allow retention of the hydrogen bonded anilino-hydrogen-diazene core, while orienting the pyridine nitrogen atoms on a trajectory favorable for coordinating metal ions. Since the low temperature emission of AzoAMoP was attributed to hydrogen bonding and from being embedded in a frozen solvent glass, we hypothesized that integrating the aAB chromophore into coordination polymers would restrict non-radiative decay pathways sufficiently to produce emissive systems at ambient temperature.

EXPERIMENTAL SECTION

General Procedures. All reagents were purchased and used without further purification. The 2,2′-diaminoazobenzene (DAAB) synthon was prepared as previously described. Toluene, dichloromethane (CH₂Cl₂), dichloroethane (DCE) and diethylether (Et₂O) were sparged with argon and dried by passing through alumina-based drying columns. All chromatography and thin-layer chromatography (TLC) were performed on silica (200–400 mesh). TLCs were developed by using CH₂Cl₂ or solvent mixtures containing CH₂Cl₂, ethyl acetate (EtOAc), hexanes, or methanol (CH₃OH). ¹H and ¹³C NMR spectra were recorded with a 500 MHz Bruker Biospin NMR instrument. Chemical shifts are reported in ppm relative to tetramethylsilane (TMS). FT-IR spectra were recorded using Bruker Optics FT-IR spectrometer equipped with a Vertex70 attenuated total reflection (ATR) accessory by collecting 1024 scans over a scan range from 4000 to 400 cm⁻¹ at 4
cm\(^{-1}\) resolution. Thermogravimetric analysis (TGA) measurements were carried out on a TA Instruments Hi-Res TGA 2950 Thermogravimetric Analyzer from room temperature to 800 °C under nitrogen atmosphere at a heating rate of 10 °C/min. LC/MS was carried on a Single Quadruple, Agilent Technologies 1200 series LC system. High resolution mass spectra were obtained at the University of Notre Dame mass spectrometry facility using microTOF instrument operating in positive ionization mode. Melting-point information was obtained using a Hydrothermal Mel-Temp instrument.

2,2′-Bis[N,N′-(2-pyridyl)methyl]diaminoazobenzene (1, AzoAMoP). AzoAMoP was synthesized as previously reported\(^4\) with minor modifications. DAAB (0.640 g, 3.02 mmol), 2-pyridinecarboxaldehyde (0.600 mL, 6.31 mmol) and 3 Å molecular sieves (0.940 g) were combined in CH\(_2\)Cl\(_2\) (30 mL) and stirred for 24 h at room temperature. Sodium triacetoxyborohydride (NaBH(OAc)\(_3\), 1.34 g, 6.32 mmol) was added, and the mixture was stirred at room temperature for 24 h. The reaction mixture was diluted with water (20 mL), and the product was extracted into CH\(_2\)Cl\(_2\) (3 × 40 mL). The combined organic layers were dried over sodium sulfate (Na\(_2\)SO\(_4\)), and the solvent was removed. Flash chromatography on silica (24:1 EtOAc/MeOH) yielded an orange-red powder (387 mg, 32.5%). Analytical data matched previously reported values.\(^4\)

2,2′-Bis[N,N′-(2-pyridyl)ethyl]diaminoazobenzene (2, AzoAEoP). AzoAEoP was synthesized as previously reported\(^5\) with minor modifications. DAAB (0.640 g, 3.02 mmol) and 2-vinylpyridine (316 μL, 2.93 mmol) were combined in CH\(_3\)OH (3 mL) and acetic acid (AcOH, 10 mL). The reaction mixture was stirred for 3 h at 45 °C before a second potion of 2-vinylpyridine (632 μL, 5.86 mmol) was added. After stirring for 24 h at 45 °C, the reaction mixture was cooled to room temperature, diluted with 10 mL of water, and the pH was adjusted to ~8 with ammonium
hydroxide (NH₄OH). The product was extracted into EtOAc (3 × 50 mL), the combined organic materials were dried over Na₂SO₄, and the solvent was removed. Flash chromatography on silica (CH₂Cl₂/CH₃OH, 25:1) gave AzoAEoP (448 mg, 35.1%) as a dark red solid. Analytical data matched previously reported values.⁵

2,2′-Bis[N,N′-(3-pyridyl)methyl]diaminoazobenzene (3, AzoAMmP). DAAB (0.640 g, 3.02 mmol), 3-pyridinecarboxaldehyde (600 μL, 6.39 mmol) and 3 Å molecular sieves (0.940 g) were combined in DCE (30 mL) and stirred for 24 h at room temperature. NaBH(OAc)₃ (1.34 g, 6.32 mmol) was added and the reaction mixture was stirred at room temperature for 24 h. The reaction mixture was diluted with water (20 mL), and the product was extracted into CH₂Cl₂ (3 × 40 mL). The combined organic layers were dried over Na₂SO₄, and the solvent was removed. Flash chromatography on silica using a gradient elution (EtOAc/hexanes 2:1→EtOAc/CH₃OH) yielded AzoAMmP as a dark red solid (532 mg, 44.7%). Diffusion of Et₂O into an CH₃CN solution of AzoAMmP provided orange-red blocks suitable for X-ray analysis. TLC Rf = 0.30 (silica, EtOAc/CH₃OH, 49:1). Mp = 158-159 °C. ¹H NMR (500 MHz, CDCl₃) δ 8.65 (s, 2 H), 8.53 (d, J = 4.9 Hz, 2 H), 7.67 (d, J = 7.8 Hz, 2 H), 7.55 (dd, J = 8.0, 1.6 Hz, 2 H), 7.25 (t, J = 6.2 Hz, 2 H), 7.19 (t, J = 7.8 Hz, 2 H), 6.76 (t, J = 7.6 Hz, 2 H), 6.70 (d, J = 8.4 Hz, 2 H), 4.50, (s, 4 H). ¹³C NMR (125 MHz, CDCl₃) δ 149.1, 149.0, 143.0, 136.6, 134.8, 134.3, 131.7, 127.3, 123.6, 116.8, 112.0, 44.7. FT-IR (neat, cm⁻¹) 3302.0, 3062.1, 3030.0, 2991.1, 2880.2, 2617.1, 2056.7, 1498.0, 1476.3, 1465.2, 1419.9, 1309.2, 1247.5, 1199.2, 1153.9, 1122.6, 1025.9, 1041.0, 906.9, 787.0, 748.0, 706.7, 601.8. HRMS (+ESI) calculated for MH⁺ 395.1979 and observed 395.1980.

2,2′-Bis[N,N′-(4-pyridyl)methyl]diaminoazobenzene (4, AzoAMPpP). DAAB (0.640 g, 3.02 mmol), 4-pyridinecarboxaldehyde (600 μL, 6.37 mmol) and 3 Å molecular sieves (0.940 g) were
combined in DCE (30 mL) and stirred for 24 h at room temperature. NaBH(OAc)₃ (1.34 g, 6.32 mmol) was added and the reaction mixture was stirred at room temperature for 24 h. The reaction mixture was diluted with water (20 mL), and the product was extracted into CH₂Cl₂ (3 × 40 mL). The combined organic layers were dried over Na₂SO₄, and the solvent was removed. Flash chromatography on silica using CH₂Cl₂/CH₃OH (10:1) yielded AzoAMP as a red orange solid (474 mg, 39.8%). Slow evaporation of chloroform (CHCl₃) provided orange-red needles suitable for X-ray analysis. TLC Rf = 0.20 (silica, CH₂Cl₂/CH₃OH, 24:1). Mp = 160-161 °C. ¹H NMR (500 MHz, CDCl₃) δ 8.58 (d, J = 5.7 Hz, 4 H), 8.55 (s, 2 H), 7.63 (dd, J = 7.9, 1.6 Hz, 2 H), 7.33 (d, J = 6.0 Hz, 4 H), 7.19 (t, J = 7.7 Hz, 2 H), 6.79 (t, J = 7.6 Hz, 2 H), 6.61 (d, J = 8.5 Hz, 2 H), 4.54 (s, 4 H). ¹³C NMR (125 MHz, CDCl₃) δ 150.2, 148.2, 143.0, 136.6, 131.8, 127.3, 122.0, 116.9, 112.1, 46.1. FT-IR (neat, cm⁻¹) 3358.0, 3111.4, 2892.3, 2130.1, 1734.2, 1652.8, 1615.2, 1528.5, 1492.4, 1415.2, 1370.5, 1312.1, 1300.9, 1244.1, 1289.1, 1206.4, 1148.1, 1124.5, 1045.8, 1016.7, 987.5, 941.5, 883.6, 804.2, 740.7, 666.3, 620.0. HRMS (+ESI) calculated for MH⁺ 395.1979 and observed 395.1985.

2,2’-Bis[N,N’-(4-pyridyl)ethyl]diaminoazobenzene (5, AzoAEP). DAAB (0.640 g, 3.02 mmol) and 4-vinylpyridine (316 μL, 2.96 mmol) were combined in CH₃OH (3 mL) and acetic acid (AcOH, 10 mL). The reaction mixture was stirred for 3 h at 45 °C before a second portion of 4-vinylpyridine (632 μL, 5.91 mmol) was added. After stirring for 24 h at 45 °C, the reaction mixture was cooled to room temperature, diluted with 10 mL of water, and the pH was adjusted to ~8 with NH₄OH. The product was extracted into EtOAc (3 × 50 mL), the combined organic materials were dried over Na₂SO₄, and the solvent was removed. Flash chromatography on silica using CH₂Cl₂/CH₃OH (15:1) yielded AzoAEP as a red orange solid (500 mg, yield 39.1%). Slow evaporation from toluene/ethanol (1:1) provided orange-red needles suitable for X-ray analysis.
TLC Rf = 0.35 (silica, DCM/CH3OH, 10:1). Mp = 149–150 °C. 1H NMR (500 MHz, CDCl3) δ 8.51 (d, J = 4.4 Hz, 4 H), 8.06 (s, 2 H), 7.32 (dd, J = 8.0, 1.6 Hz, 2 H), 7.24 (t, J = 7.8 Hz, 2 H), 7.17 (d, J = 4.7 Hz, 4 H), 6.79 (d, J = 8.4 Hz, 2 H), 6.74 (t, J = 7.5 Hz, 2 H), 3.57 (q, J = 6.4 Hz, 4 H), 2.97 (t, J = 6.9 Hz, 4 H). 13C NMR (125 MHz, CDCl3) δ 150.0, 148.1, 143.0, 136.4, 131.6, 127.2, 124.1, 116.3, 111.7, 42.9, 34.8. FT-IR (neat, cm⁻¹) 3205.4, 3046.8, 2917.4, 2851.4, 2171.9, 2330.4, 1601.0, 1565.1, 1509.1, 1458.6, 1321.6, 1204.0, 117.8, 1072.3, 1042.8, 836.7, 785.9, 368.8. HRMS (+ESI) calculated for MH⁺ 423.2292 and observed 423.2270.

{[Ag(AzoAMoP)][(CF₃SO₃)]ₙ}(AgAAMoP). AzoAMoP (25.4 mg, 64.4 μmol) in toluene (1.8 mL) was added dropwise to a toluene solution (1.8 mL) of silver trifluoromethanesulfonate (AgOTf, 16.6 mg, 64.6 μmol). Upon stirring the reaction mixture for 30 min, an orange-red solid slowly precipitated. CH₃CN (1 mL) was added to re-dissolve the precipitate, and the reaction mixture was stirred at room temperature for 2 h and filtered. Slow evaporation provided orange rectangular plates suitable for X-ray analysis. FT-IR (neat, cm⁻¹) 3361.6, 3069.5, 2904.1, 2324.2, 1981.4, 1597.2, 1581.2, 1493.6, 1466.0, 1436.7, 1371.0, 1322.3, 1285.2, 1240.9, 1219.3, 1159.1, 1107.3, 1075.2, 1052.2, 1028.3, 991.5, 888.3, 823.5, 755.2, 699.3, 633.5. Elemental analysis calcd. for C_{27}H_{25}AgF_{3}N_{7}O_{3}S (AgAAMoP·CH₃CN): C 46.79%, H 3.61%, N 14.15%; Found: C 46.50%, H 3.90%, N 14.53%. TGA shows a 0.7% weight loss between 60-120 °C, which may correspond to absorbed solvent. Decomposition occurs at 175 °C.

{[Ag(AzoAMnP)][CF₃SO₃]}ₙ(AgAAMnP). AzoAMnP (25.4 mg, 64.4 μmol) in toluene (1.8 mL) was added dropwise to a toluene solution (1.8 mL) of AgOTf (16.6 mg, 64.6 μmol). Upon stirring the reaction mixture for 30 min, an orange-red solid slowly precipitated. CH₃CN (1 mL) was added to re-dissolve the precipitate, and the reaction mixture was stirred at room temperature for 2 h and
filtered. Slow evaporation of the filtrate at room temperature provided crystals in orange-red blocks suitable for X-ray analysis. FT-IR (neat, cm\(^{-1}\)) 3230.2, 3076.0, 2941.6, 2889.8, 2362.8, 2314.7, 2165.8, 1982.2, 1862.7, 1739.6, 1603.8, 1578.9, 1507.0, 1437.0, 1431.2, 1370.4, 1297.1, 1265.5, 1254.3, 1148.0, 1024.0, 1051.3, 941.1, 741.7, 612.6, 612.3. Elemental analysis calcd. for AgAAMmP C\(_{25}\)H\(_{24.5}\)AgF\(_3\)N\(_6\)O\(_{4.5}\)S (AgAAMmP·1.5H\(_2\)O): C 44.29%, H 3.64%, N 12.40%; Found: C 44.01%, H 3.31%, N 12.02%. The TGA shows no weight loss before decomposition at 177 °C.

\([\text{Ag(AzoAMP})\text{NO}_3]_n(\text{AgAAMP})\). AzoAMP (10.0 mg, 25.4 μmol) was dissolved in 2.5 mL CH\(_3\)OH/CH\(_3\)CN (1:4) mixture with a few drops of DMF. The solution was added dropwise to an CH\(_3\)CN solution (0.5 mL) containing silver nitrate (AgNO\(_3\), 4.5 mg, 27 μmol) and tetrabutylammonium hexafluorophosphate (n-Bu\(_4\)PF\(_6\), 11.0 mg, 0.0284mmol). The reaction mixture was stirred 30 min to precipitate an orange-red solid. The solid was isolated by filtration, dissolved in 2 mL of a 1:1 mixture of CH\(_3\)OH/CH\(_3\)CN, and stirred at room temperature for 2 h. The mixture was filtered and slow evaporation provided orange-red blocks suitable for X-ray analysis. FT-IR (neat, cm\(^{-1}\)) 3516.5, 3484.4, 3262.2, 2862.0, 2826.4, 1942.1, 1639.4, 1613.5, 1610.4, 1454.6, 1454.6, 1445.0, 1322.0, 1207.8, 1206.6, 1155.8, 1100.9, 1069.1, 1028.0, 963.3, 889.5, 765.7. Elemental analysis calcd. for C\(_{24}\)H\(_{23}\)N\(_7\)O\(_3\)Ag (AgAAMP): C 51.03%, H 3.90%, N 17.37%; Found: C 50.92%, H 4.01%, N 17.43%. TGA shows a 3.6% weight loss between 60-120 °C, which may correspond to absorbed solvent. Decomposition occurs at 154 °C.

\([\text{Ag(AzoAEo})\text{NO}_3] (\text{AgAAEo})\). AzoEoP (30.0 mg, 71.1 μmol) in DCM (1 mL) was added dropwise into an CH\(_3\)CN solution (1 mL) of AgNO\(_3\) (12.0 mg, 70.6 μmol) and n-Bu\(_4\)PF\(_6\) (28.0 mg, 72.3 μmol). The reaction mixture was stirred 30 min to precipitate an orange solid. The solid was
isolated by filtration, dissolved in 2 mL of a 1:1 mixture of CH$_3$OH/CH$_3$CN, and stirred at room temperature for 2 h. The mixture was filtered and slow evaporation provided orange-red blocks suitable for X-ray analysis. FT-IR (neat, cm$^{-1}$) 3247.2, 3049.8, 2911.3, 2863.4, 2324.7, 2164.5, 2051.2, 1981.6, 1903.6, 1604.1, 1566.1, 1497.9, 1482.5, 1439.3, 1423.7, 1375.0, 1322.0, 1284.7, 1249.4, 1221.3, 1177.2, 1156.4, 1129.0, 1105.9, 1084.8, 1064.8, 1044.1, 1025.4, 1004.8, 959.6, 938.5, 882.3, 846.5, 825.4, 800.5, 762.5, 752.0, 738.4, 647.8, 616.5. Elemental analysis calcd. for C$_{26}$H$_{27}$N$_7$O$_3$Ag (AgAAEP·0.5H$_2$O): C 51.93%, H 4.53%, N 16.30%; Found: C 51.84%, H 4.43%, N 16.13%. The TGA shows no weight loss before decomposition at 166 °C.

\[[\text{Ag(AzoAEp)}_2]\text{PF}_6\]n (AgAAEP). AzoAEp (30.0 mg, 71.1 μmol) in DCM (1 mL) was added dropwise into an CH$_3$CN solution (1 mL) containing AgNO$_3$ (12.0 mg, 70.6 μmol) and n-Bu$_4$PF$_6$ (28.0 mg, 72.3 μmol). The reaction mixture was stirred 30 min to precipitate a yellow solid. The solid was isolated by filtration, dissolved in 2 mL of a 1:1 mixture of CH$_3$OH/CH$_3$CN, and stirred at room temperature for 2 h. The mixture was filtered and slow evaporation provided orange-red blocks suitable for X-ray analysis. FT-IR (neat, cm$^{-1}$) 3426.5, 3068.1, 2861.8, 2360.3, 2324.7, 2050.9, 1981.4, 1604.0, 1564.4, 1501.5, 1464.6, 1430.7, 1316.2, 1240.8, 1223.5, 1211.6, 1183.9, 1154.6, 1123.0, 1104.1, 1079.5, 1066.5, 1028.6, 939.4, 880.9, 847.3, 823.3, 800.0, 760.1, 740.8, 615.2. Elemental analysis calcd. for C$_{52}$H$_{52}$N$_{12}$F$_6$PAg (AgAAEP): C 56.89%, H 4.77%, N 15.31%; Found: C 56.72%, H 4.48%, N 15.02%. TGA shows a 0.7% weight loss between 60-120°C, which may correspond to absorbed solvent. Decomposition occurs at 197 °C.

**X-ray Crystallography.** Structural analysis was carried out in the X-Ray Crystallographic Facility at Worcester Polytechnic Institute. Crystals were glued on tip of a glass fiber or were covered in PARATONE oil on 100 μm MiTeGen polyimide micromounts and were mounted on a Bruker-
AXS APEX CCD diffractometer equipped with an LT-II low temperature device. Diffraction data were collected at room temperature or at 100(2) K using graphite monochromated Mo–Kα radiation (\(\lambda = 0.71073 \text{ Å}\)) using the omega scan technique. Empirical absorption corrections were applied using the SADABS program. The unit cells and space groups were determined using the SAINT+ program. The structures were solved by direct methods and refined by full matrix least-squares using the SHELXTL program. Refinement was based on \(F^2\) using all reflections. All non-hydrogen atoms were refined anisotropically. Hydrogen atoms on carbon atoms were all located in the difference maps and subsequently placed at idealized positions and given isotropic \(U\) values 1.2 times that of the carbon atom to which they were bonded. Hydrogen atoms bonded to oxygen atoms were located and refined with isotropic thermal parameters. Mercury 3.1 software was used to examine the molecular structure. Relevant crystallographic information is summarized in Table 1 and Table 2, and the 50% thermal ellipsoid plot is shown in Figures 1-5.

**Powder X-ray diffraction.** PXRD data were collected on a Bruker-AXS D8-Advance diffractometer using Cu-Kα radiation with X-rays generated at 40 kV and 40 mA. Bulk samples of crystals were placed in a 20 cm × 16 cm × 1 mm well in a glass sample holder, and scanned at RT from 3° to 50° (\(2\theta\)) in 0.05° steps at a scan rate of 2°/min. Simulated PXRD patterns from single crystal data were compared to PXRD patterns of experimental five AzoAXxP silver complexes, to confirm the uniformity of the crystalline samples.

**General Spectroscopic Methods.** Solution UV-Vis absorption spectra were acquired in 1.0 cm quartz cuvettes at room temperature and recorded on a Thermo Scientific Evolution 300 UV-Vis spectrometer with inbuilt Cary winUV software. Steady-state diffuse reflectance UV-Vis spectra were obtained on the same instrument with Harrick Praying Mantis diffuse reflectance accessory (Harrick Scientific Products) and referenced to magnesium sulfate. Solution emission spectra were
recorded on a Hitachi F-4500 spectrophotometer with excitation and emission slit widths of 5 nm. The excitation source was 150 W Xe arc lamp (Ushio Inc.) operating at a current rate of 5 A and equipped with photomultiplier tube with a power of 400 V.

**Emission and Quantum Yield Determination.** Steady-state emission spectra were recorded on a Hitachi F-4500 spectrophotometer with excitation and emission slit widths of 5 nm. The excitation source was 150 W Xe arc lamp (Ushio Inc.) operating at a current rate of 5 A and equipped with photomultiplier tube with a power of 400 V. Quantum yields in the solid state were determined in triplicated following published procedures using Na\(_2\)SO\(_4\) as the reference\(^{10,11}\).

**Analyte Detection by Emission.** A 100 μM suspension, with respect to AzoAE\(_p\)P units, of AgAAE\(_p\)P in toluene (2 mL) was prepared, and the emission spectra was recorded (\(\lambda_{\text{ex}} = 523\) nm). Upon the addition of each aliquot of analyte, the mixture was equilibrated by stirring for 30 min before recording the emission spectra. The emission response to all analytes was measured by integrating the emission band between 550-800 nm. Pyridine was added from a 2 mM stock solution in toluene in three equal portions, to obtain final concentrations of 33, 67, and 100 μM, and the emission was measured. After the third addition, the solvent and pyridine were removed by sparging with N\(_2\) gas, and the resulting crystalline material was re-suspended in 2 mL of toluene and stirred for 30 min before re-measuring the emission. The emission loss and restoration steps were repeated in triplicate with single analyte additions to reach 100 μM pyridine. When excess pyridine was added (500 μM), the emission completely disappeared, and could not be restored. Measurements for N-methylmorpholine (NMM) were acquired using analogous procedures, where each addition provided a final concentration of 100 μM NMM. Measurements with imidazole were acquired similarly, but emission loss was irreversible. Measurements with dimethylamine (DMA) were acquired similarly except toluene was replaced with THF, and the DMA concentration after
each switching cycle was 10 μM. After each DMA addition, the emission was restored by the addition of aliquots of 1.58 mM nitric acid (12.7 μL, 20.1 nmol) to achieve a final H⁺ concentration of 20 μM.

RESULTS AND DISCUSSION

Synthesis and structure. Using the synthetic protocols for preparing AzoAMoP and AzoAEoP, we prepared three additional aAB-bipyridyl ligands AzoAMmP (3, 2,2'-bis[N,N'-(3-pyridyl)methyl]diaminoazobenzene), AzoAMpP (4, 2,2'-bis[N,N'-(4-pyridyl)methyl]diaminoazobenzene) and AzoAEpP (5, 2,2'-bis[N,N'-(4-pyridyl)ethyl]diaminoazobenzene) from the common DAAB scaffold (2,2'-diaminoazobenzene, 6) building block (Scheme 1). Collectively, we refer to the aABs as AzoAXxP compounds where “X” represents the variable components. The m-pyridyl ligand containing the ethylene linker (AzoAEmP) has not been accessed yet owing to difficulty obtaining a stable and reactive synthon. AzoAEoP and AzoAEpP were prepared via a Michael reaction with the corresponding vinylpyridine; however, the vinyl group becomes a poor Michael acceptor with the nitrogen atom in the meta-position. The anilino nitrogen atom also appears to be a weak nucleophile as reactions with 3-(2-bromoethyl)-pyridine or 2-(pyridin-3-yl)ethyl-4-methylbenzenesulfonate failed to produce the desired product. Attempts to prepare AzoAEmP by reductive amination with 2-(pyridin-3-yl)acetaldehyde also proved unsuccessful owing to difficulty isolating sufficient quantities of the aldehyde precursor.

The five ligands can be viewed as a library of extended bipyridine ligands where each pyridine ligand can coordinate a different metal ion to form an extended network. Fortuitously, all
five AzoAXxP ligands are highly crystalline in the solid state and could be subjected to crystallographic analysis. The two pyridine nitrogen atoms of AzoAMoP are 8.37 Å apart; however, the pyridine lone pairs engage in hydrogen bonding with the anilino hydrogen atom, which orients the potential coordination sites inward. In preliminary screenings, we observed that AzoAMoP did not appear to coordinate metal ions like Zn$^{2+}$, presumably due to the thermodynamic stability of the hydrogen bonds. We reasoned that a thermodynamically stable, kinetically inert metal-ligand bond could provide access to a metal complex with AzoAMoP by shifting the equilibrium from hydrogen bonding to metal ligand bonding toward complex formation. Ag$^+$ appeared to be a good candidate for coordination polymer formation based on other successful investigations,\textsuperscript{12-16} as well as the stability of Ag–pyridine bonds.\textsuperscript{17} We further speculated that the photoactivity of Ag$^+$ might provide access to additional functionality in the resulting AB materials.

The other four AzoAXxP derivatives do not exhibit hydrogen bonding with the pyridine lone pairs, so ostensibly the ligands could bind metal ions more like typical bridging bipyridine derivatives. Two-coordinate metals with bridging bipyridine ligands are rare\textsuperscript{18-20} except for Ag$^+$.\textsuperscript{12,14,21,22} The two AzoAXxP derivatives containing methylene linkers, AzoAMmP and AzoAMPpP, can rotate at the azobenzene–anilino C–N bond, anilino–methylyene N–C bond and methylene–pyridine C–C bond. The two AzoAXxPs with ethylene linkers, AzoAEoP and AzoAEpP, can also rotate about C–C ethylene bond. The multiple degrees of freedom suggest our ligands might go through noticeable reorientation during the formation of Ag$^+$ complexes or polymers.

Although not absolutely predictive, the geometric structure of Ag$^+$ metal organic polymer chains depends on the pyridine substitution pattern, the directionality of the pyridine lone pairs,
and the conformational freedom of the ligand. Solvent and anion interactions also can impact the extended macromolecular structure. In bipyridine ligands containing ortho pyridyl nitrogen atoms, the lone pairs tend to face inward in the apo form, but often reorient during polymerization process to minimize crowding between the side chain and the Ag⁺ binding site resulting in zigzag silver chain structures. In meta- and para-pyridine bipyridine ligands, the lone pairs tend to face outward, and therefore do not necessarily need to undergo structural rearrangement in two-coordinate Ag⁺ structures. Some meta-bipyridine derivatives exhibit minimal reorientation, or asymmetric changes on one half of the molecule. In more rigid meta- and para-bipyridine ligands, linear geometries are more common because of partly or completely restricted movement.

Based on the structural trends with other bipy derivatives, we hypothesized that the pyridine nitrogen lone pairs in AzoAMoP would flip outward to accommodate Ag⁺ coordination if the intramolecular hydrogen bonds were disrupted. We prepared the AzoAMoP complex using protocols designed to gradually deliver Ag⁺ through ligand exchange with CH₃CN. A [Ag(AzoAMoP)]⁺ complex precipitated immediately from toluene solution, but re-dissolved upon addition of CH₃CN. Slow evaporation of CH₃CN led to nitrile–pyridine ligand exchange, and the formation of a AgAAMoP coordination polymer that was isolated as well-defined orange crystals. Compared to the structure of the apo ligand, the azo N–C and anilino–methylene N–C bonds in the Ag⁺ polymer with AzoAMoP ligand are rotated by 180° and 81° respectively (Figure 1). This rearrangement orients the two pyridine rings perpendicular to the azo core but retains inward-facing pyridyl nitrogen lone pairs in the formation of a helical structure where each Ag⁺ is coordinated by a pyridyl nitrogen atoms from two different AzoAMoP ligands. Ag⁺ binding increases the length of the hydrogen bonds between the anilino hydrogen atoms and the diazene
lone pairs of DAAB core by approximately 0.4 Å, and introduces a slight asymmetry in the two halves of the molecule. The core N=N bond decreases slightly from 1.28 Å to 1.26 Å owing to the decrease in hydrogen bonding (Table 1). Although the changes in hydrogen bonding might suggest a less rigid and therefore less emissive azobenzene chromophore, we expected that the metal complexation and crystal packing interactions would offset the loss of intramolecular forces.

Compared to AzoAMoP, AzoAMmP and AzoAMPp exhibit more conformational freedom, since both lack hydrogen bonding between the anilino hydrogen atoms and pyridine nitrogen atoms. The point-to-point distance between the two pyridyl nitrogen atoms are 15.96 Å and 16.52 Å respectively, but unlike a linear bridging analog like 4,4’-bipyridine, the effective distance obtained by projecting one pyridine nitrogen on a plane containing the other pyridine is shorter, 14.76 Å and 14.70 Å, respectively. We calculate the effective distance by combining the offset between the two pyridine ligands in the y and z direction where the x-axis is defined by a line going through one pyridine nitrogen atom through its para carbon atom, and defining the xy plane by that pyridine ring. The y and z offset distances are therefore the displacement from a linear bipyridine ligand (e.g. 4,4’-bipyridine) from the defined x-axis. So for AzoAMmP the pyridyl nitrogen atoms are offset from linearity by 5.41 Å (y) and 2.12 Å (z, Figure S33), and AzoAMPp has a greater displacement along the y trajectory (6.70 Å), but a shorter deviation in the z direction (1.79 Å, Figure S34). In contrast to AzoAMoP, the pyridine lone pairs in both AzoAMmP and AzoAMPp point away from the diazene core to afford extended bipyridine derivatives more reminiscent of the bridging ligands found in coordination polymers. Structural changes in meta- and para-bipy derivatives are typically minimal during Ag⁺ complex formation due to limited or no orientational freedom.22,24-26 Based on these trends, we expected the
AgAAAmP and AgAAAmP polymers, where the pyridine nitrogen atoms face outward in the apo ligands, would likewise undergo minimal structural changes.

AgAAAmP was synthesized by identical procedures used to prepare AgAAAmP. Due to limited solubility of AzoAMP ligand in toluene however, the solvent was replaced with a 1:4 mixture of CH$_3$OH/CH$_3$CN and a minimal amount of DMF, and AgOTf was substituted with AgNO$_3$. To obtain X-ray quality crystals, anion metathesis with $n$-Bu$_4$PF$_6$ was used to replace the NO$_3^-$ anion with PF$_6^-$. Compared to the structure of the apo ligand, the azo N–C bonds in the Ag$^+$ polymer with AzoAAAmP and AzoAMP ligands are rotated by 180° (Figure 2 and Figure 3), which is identical to the azo N–C bond rotation observed during the formation of AgAAAmP. This reorients the two pyridine rings, but retains outward-facing pyridyl nitrogen lone pairs in the formation of a linear or zigzag chain where each Ag$^+$ is coordinated by a pyridyl nitrogen atom from each of two different AzoAAAmP or AzoAMP ligands.

The DAAB core in AzoAAAmP and AzoAMP exhibit nearly identical the hydrogen bond lengths, 2.33 and 2.34 Å respectively; however Ag$^+$ binding produces different changes in the three derivatives containing a methylene spacer. Unlike AgAAAmP, the hydrogen bond length between the anilino hydrogen atoms and the diazene lone pairs decreases. While the hydrogen bonds in AgAMP (2.02 Å) are symmetric, AgAAAmP contains a longer (2.12 Å) and shorter (1.96 Å) bond that are both contracted compared to the apo-ligand. While AgAAAmP exhibited some asymmetry in the hydrogen bond length, the difference is much more pronounced in AgAAAmP. As expected, the increased hydrogen bonding interaction results in the opposite effect on the N=N bond lengths, which increases slightly from 1.27 Å to 1.28 Å for both coordination polymers.
The pyridine nitrogen atoms of AzoAEoP are separated by 15.04 Å. While the pyridine lone pairs face inward, the length of the ethylene linkers and ligand flexibility provide no 

*a priori*

reason to predict a reorientation would be required to coordinate Ag⁺. Using the identical procedure used to prepare AgAAMρP, a discrete monomeric complex was obtained (Figure 4). The single rotation about the azo C–N required to achieve this coordination geometry suggests kinetic trapping of a 17-membered metallacycle instead of a polymer. To the best of our knowledge, this is the largest Ag metallacycle formed from a bipydyl derivative to date. Large metallacycles with pyridyl donor groups are relatively uncommon. The most closely related examples include 13-membered-ring metallacycles of Pt and Pd,²⁷ and an 11-membered-ring metallacycle of Ag.²³

The intramolecular hydrogen bonds in AzoAEoP may be a significant factor in the preference for metallacycle formation over a polymeric structure. Two 2.36 Å hydrogen bonds between anilino hydrogen atom and the azo nitrogen atoms appear in apo ligand. When the ligand rotates to close the metallacycle, the hydrogen bonding between one of the anilino hydrogen and azo nitrogen is unavailable, and a new 2.07 Å hydrogen bond forms between the same proton and the other azo nitrogen atom. The hydrogen bonding in the upper half molecule remains unchanged, but the distance increases slightly from 2.36 to 2.38 Å. The shorter, stronger hydrogen bond formed in AgAAEoP may provide a thermodynamic driving force for metallacycle formation.

Of the five AzoAXxP ligands, AzoAEpP most closely resembles a linear bridging ligand like 4,4’-bipyridine. The point-to-point distance between the pyridine nitrogen atoms of 17.95 Å is the longest distance of all the AzoAXxP ligands, but the trajectory of the two pyridine groups is linear and the aromatic rings are coplanar in the solid state. Linear bipyridines with long nitrogen-nitrogen distances form linear Ag⁺ polymers²⁶,²⁸ and three-dimensional networks with 4-
coordinate Ag$^+$ sites.$^{29,30}$ The preference for 2-coordinate or 4-coordinate Ag$^+$ appears to correlate with the reorientation ability and rigidity of the ligands, as well as the distance between pyridine nitrogen atoms.

Although the pyridine ligands in apo-AzoAE$p$P are coplanar, rotation about the C–N bond in the ethylene spacers decreases the point-to-point distance from 17.95 Å to 6.65 Å. The effective distance is obtained similarly as described above to yield $y$ and $z$ offset distances of 2.22 and 9.09 Å, respectively (Figure S35). Each Ag$^+$ is four-coordinate with each pyridine coming from one of four unique AzoAE$p$P ligands. We partially attribute the increased coordination to the reduced steric requirements at the donor ligand site of the para-pyridine isomer. Each AzoAE$p$P ligand also coordinates two separate Ag cations to form a three-dimensional polymer with the distorted tetrahedral Ag$^+$ sites (Figure 5B). The lack of steric hindrance at the coordination site of the para-pyridine ligands appears to be an important prerequisite for three-dimensional polymer formation with bipyridines. In previous Ag$^+$ polymers and three-dimensional networks with 4-coordinate Ag$^+$ sites, most of the ligands undergo minimum or no reorientation.$^{26,28,30}$ Unlike these linear bipyridine ligands, AzoAE$p$P exhibits a high degree of freedom, which allows optimization of steric and electrostatic factors in Ag$^+$ complex formation.

**Emission.** AzoAMoP exhibits no measurable emission at room temperature, but emits when frozen in a solvent glass at 77 K. The four additional AB compounds behave similarly. Although the wavelengths vary, all the five compounds emit with a $\lambda_{\text{max}}$ between 566 nm and 610 nm (Table 2). Similar to AzoAMoP and AzoAEoP,$^{4,5}$ none of the new AzoAXxP derivatives exhibit significant evidence for photoisomerization upon irradiation. While quantitative measurements of minimally emissive complexes are imprecise, qualitatively, AzoAEoP and AzoAE$p$P exhibit
brighter emission than AzoAM\textsubscript{m}P and AzoAM\textsubscript{p}P. AzoAM\textsubscript{o}P appears to emit more weakly than the other four derivatives.

To assess the photochemistry of the Ag\textsuperscript{+} complexes, the emission of AgAAM\textsubscript{o}P, AgAAM\textsubscript{m}P, AgAAM\textsubscript{p}P, and AgAAE\textsubscript{p}P were evaluated in toluene as suspensions of powdered crystals. The four coordination polymers form semi-homogeneous dispersions in toluene that remain suspended for several months. Similar to the free ligands, the emission $\lambda_{\text{max}}$ of the Ag polymers occurs around 600 nm for all the complexes (Table 2). In the solid state, the relative integrated emission intensities reveal a nearly 30-fold brighter emission for the four-coordinate AgAAE\textsubscript{p}P complex compared to the least emissive AgAAM\textsubscript{p}P complex (Figure 6). Complex formation redshifts the absorbance maximum of AzoAE\textsubscript{p}P from 502 nm to 594 nm with an emission peak centered at 641 nm (Figure 7). Similar to the absorbance, the emission of AgAAE\textsubscript{p}P redshifts 31 nm from that of free AzoAE\textsubscript{p}P measured at 77 K, which is a greater a change than the other three polymeric complexes. The luminescence response appears to validate the initial hypothesis that embedding the AzoAX\textsubscript{x}P ligand in a solid state material increases the degree of radiative decay of the AB excited state.

The emission of AB in solution at room temperature is too weak to be detected by conventional emission instruments owing to efficient decay of the excited state exist via non-radiative pathways.\textsuperscript{31} In many aromatic chromophores, the emission of a photon accompanies a $S_1(\pi\pi^*)\rightarrow S_0$ transition as the excited molecule returns to the ground state. In AB however, the $S_2(\pi\pi^*)\leftrightarrow S_0$ absorbance has the largest extinction coefficient, but a weakly absorbing interstitial $S_1(n\pi^*)$ state is present. Although the photophysical processes remain an active area of investigation,\textsuperscript{32-34} a rapid intersystem crossing to a vibrationally excited $S_1(n\pi^*)$ state occurs following $S_2(\pi\pi^*)\leftrightarrow S_0$ absorbance. AB isomerization then occurs via the concerted inversion
pathway from the S$_1$ state.$^{35,36}$ A modest increase in emission intensity can be observed in a frozen matrix when relaxation pathways involving molecular motion are impaired.

Radiative decay of AB can be increased by manipulating the frontier orbitals of the chromophore as observed in 2-borylazobenzenes.$^{37-40}$ Engaging the diazene lone pair in a dative bond with a boron atom lowers the energy of the lone pair, or B–N bonding orbital, below the diazene π–bond. Thus forming the nπ* state would require promoting an electron into the half-occupied, higher energy diazene π-orbital. The absence of the interstitial S$_1$(nπ*) state removes the optically forbidden S$_1$(nπ*)→S$_0$ transition and opens the S$_2$(ππ*)→S$_0$ emission channel.

We hypothesized that introducing intramolecular hydrogen bonds between the diazene lone pair and anilino hydrogen atom might lead to a similar, albeit less drastic, change in the frontier orbitals to those observed in 2-borylazobenzenes, increase the radiative decay of the excited state. This hypothesis was supported by our previous observations that an AzoAMoP derivative lacking intramolecular hydrogen bonds had significantly weaker emission at 77 K similar to the luminescence of AB under the same conditions.$^4$ To support the increased emission hypothesis, we interrogated the ground state electronic structure of each ligand using Density functional theory (DFT). In order to capture the structural impact of the solid-state environment on the electronic structure for each ligand, unit cell vectors and atom positions were determined using three-dimensional periodic boundary conditions before extracting a single molecule for analysing the electronic structure, an approach we have previously applied to luminescence metal-organic systems.$^{41}$ The first ten excited states were computed vertically using time-dependent DFT (TDDFT) for all the ligands.
The computational approach provides values that correlate closely with experimentally measured absorption energies. As expected, the gap between the n and π* orbitals decreases significantly going from AB to DAAB, but the order remains unchanged (Figure S36). The red-shifting of the ππ* absorption maxima between AB and DAAB was calculated to be 0.95 eV, in reasonable agreement with the experimental value of 1.23 eV. Likewise, the calculated and experimental values for the five aAB showed reasonable agreement. The minimum and maximum red-shifting of the ππ* transition of the five aAB ligands were measured as 0.139 eV and 0.142 eV with respect to DAAB. The equivalent red-shifts determined computationally were 0.250 eV and 0.294 eV, a similar spread to the experimental values shifted by approximately 0.1 eV.

Since our series of modified aAB ligands are significantly more emissive in a frozen matrix than AB and DAAB, we reasoned that the anilino substituent must induce further changes in the frontier orbitals. In examining the frontier orbital energies, we observed that the pyridine moieties induce a stabilization of the diazene-centered n-type orbital for the four aAB ligands (maximum 0.33 eV). This observation is in line with the expectation that the inductive effect of the pyridine substituent results in a slightly more electron rich secondary anilino nitrogen atom, leading to a slightly more stable π* orbital and making the diazene lone pairs somewhat better hydrogen bond acceptors. AzoAMoP, which exhibits a relative orbital destabilization of approximately 0.44 eV, provides an exception to this trend. AzoAMoP shows a reduced gap between the n and π* orbitals compared to the other four derivatives. We attribute this difference to the additional hydrogen bonding provided by the pyridine substituent, reducing the interaction between the diazene lone pair and the anilino hydrogen atom. Introducing a meta/para nitrogen atom within the pyridine unit or an ethylene spacer increases the distance between the nitrogen donor atom and the diazene core rendering this interaction negligible and yielding a net stabilization of the lone pair, as seen
in the other four derivatives. The lack of emission in solution and subsequent increase in a frozen matrix is also consistent with hydrogen bonding as a key contributor to the emission behavior. The restricted motion at low temperature would be expected to lead to a stabilization of the hydrogen bonds and therefore the energy levels of the frontier orbitals, which would not necessarily occur in solution.

The electron donating pyridine moieties destabilize the $\pi$ (HOMO) orbital for all ligands with respect to the DAAB control molecule. This destabilization exceeds the stabilization of the diazene lone pair orbital (LUMO), which results from hydrogen bonding between the diazene lone pair and the anilino hydrogen atom (maximum 0.95 eV). AzoAMoP again provides an exception, where the $\pi^*$ orbitals are stabilized upon the addition of the pyridine substituents (0.52 eV). The combination of these effects results in a general stabilization of the $\pi\pi^*$ excited state and destabilization of the $n\pi^*$ state, causing an inversion of the emissive $\pi\pi^*$ and non-emissive $n\pi^*$ states with respect to both AB and DAAB. As expected, this results in the removal of the lower-lying $n\pi^*$ state as a radiationless decay channel.\(^{40}\) Notably, the inversion of states is the least pronounced in AzoAMoP. In an absolute sense, the $n\pi^*$ state still lies higher in energy than the $\pi\pi^*$, but the small energy gap (0.04 eV) could permit internal conversion between these two states, accounting for the lower emission intensity of AzoAMoP.

To generate emission at room temperature, we hypothesized that embedding the ligands in coordination polymers would provide the requisite restrictions on molecular motion needed to stabilize the diazene lone pair-anilino hydrogen atom hydrogen bond. Our hypothesis was validated by the emission measurements on the Ag\(^+\) systems. The 2.43 Å diazene lone pair-anilino hydrogen atom bond length in [Ag(AzoAMoP)]\(_n\) is the longest of the four polymeric complexes, and correlates with the lowest relative integrated emission intensity (Table 2). As the hydrogen
bond length shortens in $[\text{Ag(AzoAMmP)}]_n$ (2.12 Å) and $[\text{Ag(AzoAMP)}]_n$ (2.03 Å), the relative emission intensity increases. As our hypothesis predicts, $[\text{Ag(AzoAEpP)}]_2)_n$ has the brightest emission and the shortest hydrogen bonds. In order to understand the effects of the structural reorganization imposed by $\text{Ag}^+$ coordination on the frontier orbitals and excited states, the same computation procedures were applied to the polymeric complexes. These calculations only capture the structural effects of the $\text{Ag}^+$ coordination since a neutral ligand is extracted from the periodic, solid state calculations; therefore, the model does not account for the electronic effects of $\text{Ag}^+$ coordination.

While the trend between hydrogen bond length and emission intensity correlates as expected, the computational results predict a small energy gap between the n and $\pi^*$ orbitals. Since the structural reorganization of the anilino moieties after polymer formation causes only a slight modulation in the lowest-lying excited states, the qualitative ordering of the states remains unchanged with respect to the results obtained for the crystalline ligands alone. Interestingly, $[\text{Ag(AzoAEpP)}]_2)_n$ is over an order of magnitude more emissive than the other three polymeric complexes, which indicates other contributing factors to the photophysical behavior. The calculations based on extracted ligands only capture structural effects imposed by coordination to $\text{Ag}^+$. While we are able to demonstrate a correlation between hydrogen bond length and emission behavior across the series of materials, we cannot draw any conclusions regarding the influence of $\text{Ag}^+$ atoms or the crystalline environment on orbital or excited state energies. Together, these results show that the enhanced luminescence observed in these azobenzene derivatives could be based on the inversion of the aforementioned electronic excited states relative to azobenzene, but no conclusions can be made concerning the differences in emission intensity observed upon the
formation of silver complexes, or the differences in luminescence intensity observed between different silver-based polymers.

AgAAEₚP exhibits the brightest emission of all the Ag⁺ coordination polymers with a quantum yield of 0.8%. The other complexes have quantum yields of less than 0.1%, and therefore the exact values are not reliable. We examined the emission response to the Ag⁺-coordinating analytes pyridine, N-methylmorpholine, DMA and imidazole. When 5 equivalents of pyridine with respect to AzoAEₚP units is added to a suspension of AgAAEₚP in toluene, no emission was detected; however, the absorbance spectrum showed the presence of AzoAEₚP. When a sub-stoichiometric amount of pyridine was added, the coordination polymer emission decreased and the absorbance spectrum showed evidence for the release of some AzoAEₚP (Scheme 2). We suspected that pyridine was displacing AzoAEₚP pyridine ligands bound to the Ag⁺ sites, which would cause the loss of extended structure in the coordination polymer. The release of AzoAEₚP, which is not emissive at room temperature in solution, would produce an on/off switching behavior. To provide supporting evidence for this signaling mechanism, the pyridine was removed by sparging the solution with N₂ gas. The recovery of emission after the evaporation and re-dispersion of the material in toluene suggests the reassembly of the original AgAAEₚP structure. The disassembly/reassembly process can also be observed by PXRD. Upon the addition of pyridine, the diffraction pattern indicates the formation of amorphous material, and the reassembled material exhibits an identical spectrum to the original coordination polymer.

The reversible structure disassembly process can be monitored by emission spectroscopy. Initially, the emission spectrum of AgAAEₚP exhibits features with maxima at 620 nm (Figure 8A). Upon the addition of pyridine, the peak intensity decreases gradually, and correlates with expected changes in absorbance from AzoAEₚP. The pyridine removal process restores the
original spectrum characteristic of AgAAEP. This process is fairly robust, and can be repeated with minimal loss of emission. After five cycles, the material retains 90% of the original emission intensity (Figure 8B), and the slight erosion of the maximum response is likely due to irreversible loss of structure, or incomplete removal of pyridine. NMM shows similar results to pyridine. The boiling points of pyridine and NMM are 115.2 °C and 116 °C, respectively, so the sparging with N\textsubscript{2} removes both the analyte and solvent (toluene, bp 111°C).

In contrast, when exposed to imidazole, the emission loss is not reversible. Although imidazole displaces AzoAE\textsubscript{P} from the coordination sphere of Ag\textsuperscript{+} in the coordination polymer, the lack of volatility (bp 256 °C) makes removal nearly impossible. A similar response occurs upon treatment of AgAAEP with potassium bromide. The removal of Ag\textsuperscript{+} through the formation of insoluble halide complexes leads to an irreversible loss of emission and a UV spectrum consistent with free AzoAE\textsubscript{P}.

The ability to reversibly detect analytes through changes in extended structure is not limited to removal by evaporation. Addition of DMA leads to a loss of emission analogous to the response to pyridine. The addition of dilute nitric acid causes the AgAAEP to reassemble. This process can also be repeated for multiple cycles without significant loss of absolute emission intensity (Figure 8C). The emission loss and restoration process by adding and removing volatile analytes like pyridine and NMM demonstrates the viability of a disassembly-reassembly process for emission detection. While physical/electrostatic interactions are more common signal transduction pathways,\textsuperscript{42-44} our system provides an alternative sensing mechanism driven by coordination events, but primarily demonstrates that the coordination polymer is required for the AzoAX\textsubscript{xP} ligands to be emissive at room temperature.
CONCLUSION

Although ABs are not typically emissive, embedding AB chromophores in a rigid coordination polymer can enhance the degree of radiative decay of the excited state. In the aAB ligands investigated, both Ag⁺ coordination and intramolecular hydrogen bonds contribute to the emission enhancement; however, the hydrogen bonding accounts for most of the changes in the frontier molecular orbitals. The stabilization of the ππ* excited state below the interstitial nπ* usually involved in the non-radiative decay of the excited state leads to luminescence in both solid state and crystalline samples dispersed in solvent. Our initial investigations suggest three-dimensional networks are more emissive than linear coordination polymers, however, additional examples will be required to confirm this hypothesis. The coordination polymers can be completely, or partially disassembled by the addition of Ag⁺-binding analytes such as amines, and reformed after analyte removal to provide a sensor-like system. This signal transduction mechanism differs from many polymeric sensors, and will be investigated as an alternative approach to designing practical luminescent probes.
**Figure 1A.** Preparation of AgAAMoP showing the thermal ellipsoid representation of AzoAMoP and AgAAMoP at the 50% probability level and selected atom labels. Hydrogen atoms except for those engaged in intramolecular hydrogen bonds are omitted for clarity. The triflate anion and a non-coordinating CH$_3$CN solvent molecule are omitted for clarity. AgAAMoP was prepared in toluene using AgOTf as the silver source using a slow ligand exchange process with CH$_3$CN. During the polymer formation, the azo N–C and anilino–methylene N–C bonds rotate by 180° and 81°, respectively. **1B.** Thermal ellipsoid representation at the 50% probability level of the expanded helical polymer chain. Hydrogen atoms, triflate anions and CH$_3$CN solvent molecules are omitted for clarity.
Figure 2A. Preparation of AgAAMmP showing the thermal ellipsoid representation of AzoAMmP and AgAAMmP at the 50% probability level and selected atom labels. Hydrogen atoms except for those engaged in intramolecular hydrogen bonds are omitted for clarity. The triflate anion is removed for clarity. AgAAMmP was prepared in toluene using AgOTf as the silver source using a slow ligand exchange process with CH$_3$CN. During the polymer formation, the azo N–C bonds rotate by 180°. 2B. Thermal ellipsoid representation diagram at the 50% probability level of the expanded helical polymer chain. Hydrogen atoms, triflate anions are omitted for clarity.
**Figure 3A.** Preparation of AgAAMpP showing the thermal ellipsoid representation of AzoAMpP and AgAAMpP at the 50% probability level and selected atom labels. Hydrogen atoms except for those engaged in intramolecular hydrogen bonds are omitted for clarity. The PF$_6$ anion is removed for clarity. AgAAMpP was prepared in CH$_3$OH/CH$_3$CN (1:4) using AgNO$_3$ as the silver source. Subsequent anion exchange with n-Bu$_4$PF$_6$ in CH$_3$CN provided x-ray quality crystals. During the polymer formation, the azo N–C bonds rotate by 180°. **3B.** Thermal ellipsoid representation showing at the 50% probability level of the expanded zigzag polymer chain. Hydrogen atoms, PF$_6$ anions are omitted for clarity.
Figure 4. Preparation of AgAAEoP showing the thermal ellipsoid representation of AzoAEoP and AgAAEoP at the 50% probability level and selected atom labels. Hydrogen atoms except for those engaged in intramolecular hydrogen bonds are omitted for clarity. The PF$_6$ anion is removed for clarity. AgAAEoP was prepared in CH$_3$OH/CH$_3$CN (1:4) using AgNO$_3$ as the silver source with added $n$-Bu$_4$PF$_6$ to provide x-ray quality crystals. During the complex formation, one azo N–C bond rotates by 180˚ while the other remains stationary to form a 17-membered ring. The two hydrogen bonds formed between two anilino hydrogen atoms and the same azo N atom are asymmetric with lengths of 2.07 Å and 2.38 Å respectively.
Figure 5A. Preparation of AgAAEpP showing the thermal ellipsoid representation of AzoAEpP and AgAAEpP at the 50% probability level and selected atom labels. Hydrogen atoms except for those engaged in intramolecular hydrogen bonds are omitted for clarity. The PF$_6$ anion is removed for clarity. AgAAEpP was prepared in DCM using AgNO$_3$ as the silver source with added $n$-Bu$_4$PF$_6$. The resulting precipitate was dissolved in CH$_3$CN to yield x-ray quality crystals from slow solvent evaporation. During the polymer formation, the ethylene C–C bonds rotate by 180° and ethylene-pyridine C–C rotates minimally. 5B. Thermal ellipsoid representation at the 50% probability level of the three dimensional polymer chain. Hydrogen atoms and PF$_6$ anions are omitted for clarity.
Figure 6. Solid state emission spectrum of the five AzoAXxP silver complexes ($\lambda_{ex}$ 523 nm) showing the relative intensities.
Figure 7. Solid state diffuse reflectance (black) and solid state emission ($\lambda_{ex} = 523$ nm, pink) spectra of AgAAEP. Inset: cuvettes containing a suspension of the complex crystals in toluene (100 $\mu$M) without irradiation (left), and excited with 365 nm light (right).
Figure 8A. Normalized emission response of AgAAEpP to pyridine. AgAAEpP (100 μM) was suspended in toluene (black), treated with 100 μM pyridine (red), and redispersed in toluene after removing the solvent and analyte by sparging with N₂ (blue). The response also shows stepwise decreases when incremental amounts of pyridine were added to reach final concentrations of 33 μM, 67 μM, and 100 μM. (inset). 8B. Normalized emission response of AgAAEpP to multiple
cycles of pyridine addition and removal. AgAEpP (100 μM) was exposed to 100 μM of pyridine followed by a sparging and redispersal process a total of 4 times. Normalized emission response of AgAEpP to multiple cycles of DMA addition and removal. AgAEpP (100 μM) was exposed to 10 μM of DMA followed by an addition of nitric acid to reach a final concentration of 20 μM H⁺ a total of 4 times.
Scheme 1. Synthetic protocols for preparing five AzoAXxP ligands. Reagents and conditions: (a) NaBH(OAc)$_3$, CH$_2$Cl$_2$, room temperature; (b) AcOH, MeOH, 45 °C.
Scheme 2. Proposed action of the on–off response of AgAAEₚP. Addition of Ag coordinating ligands like pyridine leads to loss of the polymeric structure necessary to induce luminescence from the azobenzene chromophore. Removal of pyridine leads to a reassembly of the coordination polymer.
Table 1. Selected Interatomic Distances (Å) and Angles (deg) for the five AzoAXxP Ligands.

|          | Selected bond lengths                     | Bond angles                                      |
|----------|-------------------------------------------|-------------------------------------------------|
| AzoAMoP  | N(3)–N(3) 1.2793(15)                      | N(3)–N(3)–C(12) 116.87(10)                     |
|          | N(3)–C(12) 1.4127(16)                     | N(3)–C(12)–C(7) 126.89(11)                     |
|          | C(12)–N(2) 1.3573(18)                     | N(2)–C(7)–C(12) 121.64(11)                     |
|          | N(2)–C(6) 1.4434(18)                      | C(6)–N(2)–C(7) 123.93(11)                      |
|          |                                          | N(2)–C(6)–C(5) 110.70(10)                      |
|          |                                          | N(1)–C(5)–C(6) 118.15(12)                      |
| AzoAMmP  | N2–N2 1.268(3)                             | N2–N2–C2 114.9(2)                               |
|          | C2–N2 1.414(3)                             | N2–C2–C3 115.07(17)                             |
|          | C3–N1 1.373(3)                             | N1–C3–C2 119.75(18)                             |
|          | C8–N1 1.444(3)                             | C3–N1–C8 122.48(17)                             |
|          |                                          | N1–C8–C4 113.39(17)                             |
| AzoAMPp  | N1–N1 1.265(3)                             | N1–N1–C7 115.08(18)                             |
|          | C7–N1 1.415(2)                             | N1–C7–C12 115.58(15)                            |
|          | C12–N2 1.373(2)                            | N2–C12–C7 119.91(15)                            |
|          | C1–N2 1.449(2)                             | C12–N2–C1 122.46(15)                            |
|          |                                          | N2–C1–C2 115.93(16)                             |
| AzoAEoP  | N1–N1 1.267(3)                             | N1–N1–C1 115.48(18)                             |
|          | C1–N1 1.408(2)                             | N1–C1–C6 115.84(15)                             |
|          | C6–N2 1.354(2)                             | N2–C6–C1 120.30(16)                             |
|          | C7–N2 1.446(2)                             | C6–N2–C7 124.91(17)                             |
|          | C9–N3 1.340(2)                             | N2–C7–C8 112.68(15)                             |
|          |                                          | N3–C9–C8 117.21(17)                             |
| Compound | H-bond distances NH···N=N(Å) | ε [M⁻¹cm⁻¹] | λ_max (nm) | Em_max (nm) | Relative emission intensity (RT) |
|----------|-----------------------------|-------------|-----------|-------------|-------------------------------|
| AzoAMoP  | 2.046(15)                   | 12334       | 495       | 602^[a]     | n.a.                          |
| AzoAMmP  | 2.33(3)                     | 13729       | 497       | 607^[a]     | n.a.                          |
| AzoAMpP  | 2.34(2)                     | 13702       | 495       | 603^[a]     | n.a.                          |
| AzoAEoP  | 2.36(2)                     | 15874       | 501       | 619^[a]     | n.a.                          |
| AzoAEpP  | 2.00(3)                     | 15195       | 502       | 616^[a]     | n.a.                          |
| AgAAMoP  | 2.43(3), 2.47(5)            | n.a.        | 440       | 600^[b]     | 0.035                         |
| AgAAMmP  | 2.12(9), 1.96(2)            | n.a.        | 440       | 618^[b]     | 0.067                         |
| AgAAMpP  | 2.02(3)                     | n.a.        | 475       | 590^[b]     | 0.069                         |
| AgAAEpP  | 2.00(3), 2.05(1), 2.01(3), 2.02(2) | n.a. | 508       | 641^[b]     | 1.0                           |

[a] λ_ex = 490 nm, [b] λ_ex = 523 nm

Table 2. Photophysical Properties of AzoAXxP Ligands and Coordination Polymers.
Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: XXXXXXXXX.

- $^1$H and $^{13}$C NMR spectra for new AzoAXxP compounds synthesized. TGA and PXRD spectra for all new Ag$^+$ coordination polymers and complexes reported.

- Room temperature and 77K emission spectra of all AzoAXxP compounds. Diffuse reflectance and emission spectra of all Ag$^+$ coordination polymers. Emission data for analyte additions to AgAAEP.$^+$

- Diagrams of Ag$^+$–AzoAXxP units showing distance calculations based on the trajectory and positions of the pyridine units.

- Additional computation details, figures and tables of calculated energies and bond lengths

- Complete X-ray tables and fully labeled thermal ellipsoid representations of all coordination polymers and complexes.

Accession Codes

CCDC 1854329–1854336 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.
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SYNOPSIS

Four luminescent silver coordination polymers have been assembled using unique bipyridine ligands containing an azobenzene chromophore. The most emissive complex, AgAAEP, containing tetra-coordinate silver sites, and undergoes reversible disassembly with other silver binding ligands to create a luminescent switch.