A New Standard in High-Field Terahertz Generation: the Organic Nonlinear Optical Crystal PNPA


ABSTRACT: We report the full characterization of a new organic nonlinear optical (NLO) crystal for intense THz generation: PNPA ((E)-4-((4-nitrobenzylidene)amino)-N-phenyl-aniline). We discuss crystal growth and structural characteristics. We present the wavelength dependence of THz generation, the thickness dependence of the THz spectrum for PNPA crystals and measure the efficiency. PNPA enables intense THz generation that surpasses NLO crystals DAST and OH-1, which have been the standard in organic high-field THz generators for several years. With our experimental conditions, PNPA can generate a peak-to-peak field strength of 2.9 MV/cm compared to 2.2 and 1.5 MV/cm from DAST and OH-1, respectively. This corresponds to a THz generation efficiency exceeding 4%.

KEYWORDS: terahertz generation, nonlinear optical crystal, high-field terahertz

INTRODUCTION

The use of intense, broadband pulses of terahertz (THz) light as an ultrafast excitation source is enabling cutting-edge condensed-phase measurements. The relatively low-frequency THz light can resonantly excite collective modes, which efficiently drives motion far from equilibrium conditions. This efficient resonant excitation has enabled the direct measurement of anharmonic potential energy surfaces, the direct excitation and measurement of an electromagnon in multiferroic TbMnO₃, and the transition to a metastable ferroelectric state in SrTiO₃. Experimental setups with the capability to generate a pair of intense THz pulses can also facilitate 2D THz spectroscopic measurements and have provided additional insights on a variety of fascinating systems.1−10

Each of these experiments require high-field THz generation, and intense THz pulses were provided by organic nonlinear optical crystals like OH-1,1,2 DSTMS,2,4−6,10 or DAST11 or by inorganic LiNbO₃ in a tilted-pulse-front configuration.4,6 DAST, the original organic crystal for high-field THz generation, was developed as a nonlinear optical crystal over 30 years ago.11,12 Ten years ago, DAST was shown to be extremely efficient with THz generation, demonstrating peak field strengths in excess of 1 MV/cm.13 Since that demonstration, several new THz generators have been developed, showing marginal improvement in THz output.16−23 However, no new organic THz generator has significantly outperformed DAST.

DAST and similar organic crystals have been the standards in THz generation because they generate field strengths exceeding 1 MV/cm, they boast generation efficiencies greater than 1%, and they can produce broad bandwidths that are commensurate with the pump pulse duration. These crystals boast these capabilities due to a fast electronic NLO response and relatively large NLO coefficients. Because of the exciting experiments that high-field THz sources have enabled, this begs the question of what makes an ideal THz source and if anything can replace the standard that these organic crystals have set.

Ideal THz generators would create high peak-field strengths and smooth spectra without absorption dips. Despite their large NLO coefficients, DAST and OH-1 exhibit strong absorptions (>200 cm⁻¹) and imperfect phase matching leading to absorption dips at 1 (DAST), 3, and 4.5 THz (OH-1). A new standard would ideally have larger NLO coefficients enabling generation efficiencies above 3% and absorption coefficients less than 200 cm⁻¹ across a broad range of THz frequencies.

Our laboratories recently disclosed a new data mining approach to discover and produce organic THz crystals.24 We mined the Cambridge Structural Database (CSD) and combined the structural information with first-principles calculations to identify new organic materials with ideal molecular and solid-state properties for THz generation. Of these new materials, PNPA ((E)-4-((4-nitrobenzylidene)-...
amino)-N-phenylaniline) was demonstrated to output the highest intensity THz generation, rivaling that generated by DAST and OH-1, necessitating a full characterization to showcase its THz generation capabilities. In this report, we provide an in-depth analysis of the synthesis, growth, crystal characterization, and optical properties of PNPA and demonstrate that it enables THz generation that surpasses current state-of-the-art NLO crystals DAST and OH-1. Because PNPA demonstrates the capability to outperform current standards by generating significantly stronger field strengths with an extremely high generation efficiency, we suggest that PNPA should be used as a new standard in high-field THz generation.

**EXPERIMENTAL SECTION**

**Synthesis of PNPA.** Equimolar amounts of 4-nitrobenzaldehyde (15.1 g, 0.1 mol) and 4-aminodiphenylamine (18.4 g, 0.1 mol) were added to a 500 mL flask. Next, 300 mL of ethanol and 3.0 g of anhydrous sodium sulfate were added. The reaction was stirred under an argon atmosphere at 80 °C for 1 h. The ethanol was removed by vacuum evaporation (rotovap), and 300 mL of ethanol was again added to the reaction. The mixture was again stirred for 1 h at 80 °C, followed by removal of solvent. The solid product was then dissolved in 1 L of CH₃Cl₂, and the solution was filtered through a silica gel column using CH₂Cl₂ as eluent to provide the desired product as a solid (31.7 g, 100% yield). M.P. = 176.5 °C. ¹H NMR (500 MHz, CDCl₃), δ 8.61 (s, 1H), 8.33–8.31 (d, J = 8.8, 2H), 8.07–8.05 (d, J = 8.8, 2H), 7.33–7.30 (m, 4H), 7.12–7.11 (m, 4H), 7.00–6.98 (t, J = 7.4, 1H), 5.86 (s, 1H).

**Large Crystal Growth.** Previous studies have reported that large PNPA crystals can be grown from chloroform and methanol mixtures via slow evaporation. Our additional studies demonstrate that crystal growth via slow evaporation from CH₃Cl₂ provides the best quality crystals with the largest sizes. In a typical slow evaporation experiment, 3.3 g of powdered (pure) PNPA was dissolved in 100 mL of CH₂Cl₂ with heating and stirring until all solids are dissolved. The resulting solution was allowed to slowly evaporate at ∼1.5 mL per day at room temperature over the course of 2 weeks. During this time, red, prismatic seed crystals began to form. The resulting seed crystals were extracted after the 2 weeks and resubjected to a solution concentrated at 3.1 g of PNPA per 100 mL of CH₂Cl₂ at the same rate and for the same amount of time. Resulting crystals recovered from solution ranged in size from 5 mm × 10 mm to 10 mm × 10 mm with varying thicknesses due to the prismatic shape of the crystals. Under the same conditions, some flat crystals were grown with uniform thickness. High quality crystal windows of varying thickness were then cleaved from the prisms and polished to desired thicknesses. Figure 1 shows examples of (a) prismatic as-grown PNPA crystals and (b) a cleaved and polished crystal.

![Figure 1](https://example.com/figure1.png)

**Structure Confirmation and Face Determination.** Single-crystal X-ray diffraction was used to confirm that our PNPA grown under the above method maintained the same crystal structure as the reported structure in the database (CCDC ID: 991743). We also identified the main crystal face for irradiation (010) using powder X-ray diffraction (see section 2 in the Supporting Information for experimental details). XRD measurements show that no other compound had cocrystallized with PNPA, indicating that the grown compound was high-purity PNPA. NMR measurements confirmed a >97% purity.

**Terahertz Generation Measurements.** Measurements were performed using ultrashort pump pulses with a broad range of NIR wavelengths. To generate these wavelengths, the 800 nm light generated from a Ti:sapphire regenerative amplifier is sent into an optical parametric amplifier which outputs ∼100 fs light pulses across the whole range from 1250 to 2150 nm (1250–1600 nm signal, 1600–2150 nm idler). Specific pump parameters for each measurement are listed in conjunction with each figure, and a summary is given in the Supporting Information in Table S2. As shown in Figure 2, the signal or idler pump pulses are directed through a chopper, which reduced the repetition rate from 1 kHz to 500 Hz, and to the THz generation crystal where excess pump light is removed using a 2 mm thick HDPE filter. The transmitted THz is expanded using a 1:5 telescope before being focused down to the sample position. EO detection is used to record traces. The THz pulses are then directed through a 1:5 telescope before being focused down to the sample position. EO detection is used to record traces.

![Figure 2](https://example.com/figure2.png)

**RESULTS AND DISCUSSION**

**Molecular Property and X-ray Crystal Structure.** Simple models show that three of the key factors that govern the THz generation efficiency in a NLO crystal include the molecular hyperpolarizability, molecular alignment in the crystalline state,
and the irradiated crystal face. Through our initial data mining effort, the hyperpolarizability of PNPAmolecules was calculated to be $316 \times 10^{-30} \text{esu}$, which is significantly larger than the values for OH-1 ($89.6 \times 10^{-30} \text{esu}$) and DAST ($201 \times 10^{-30} \text{esu}$).

In order to quantify the molecular alignment, the crystal packing order parameter ($\cos^3(\theta_p)$) was calculated, where $\theta_p$ is defined as the angle between the molecular hyperpolarizability vector and the polar axis of the unit cell. An order parameter of 1 indicates an ideal head-to-tail molecular alignment and a value of 0 indicates centrosymmetric packing without NLO activity. The value of $\theta_p$ for PNPAm is 0.66°, giving an order parameter that is 0.9998 (see Figure 3). Compared to the order parameter values of OH-1 (0.69) and DAST (0.83), PNPAm has a nearly ideal molecular alignment for THz generation.

**THz Generation and Optical Characterization.** The nearly noncentrosymmetric packing and high hyperpolarizability of PNPAmake it a strong candidate for high intensity THz generation, which we predicted to surpass both DAST and OH-1. Here we provide a comprehensive description of the THz generation capabilities of PNPAm and we directly compare PNPAm to current standard THz generators DAST and OH-1.

PNPAm generates a strong, single-cycle pulse with the capability to reach peak-to-peak electric-field strengths >2 MV/cm, as shown in Figure 4a. Figure 4b shows the associated spectrum with an absorption dip at 2 THz and frequencies extending out to 5 THz. As shown in Figure 4b, as well as Figures 5, 6, and 8, PNPAm generates a relatively broad and smooth spectrum due to the THz absorption coefficient being <200 cm$^{-1}$, even for the largest absorption at 2 THz (see Supporting Information). We also tested multiple crystals to ensure that crystal growth was replicable and led to consistent THz output. The insets of Figure 4 show the THz traces and spectra of 10 PNPAm crystals, with thicknesses ranging from ∼300 to 900 μm. Small variabilities arise in spectral shape from differences in crystal thickness and quality. In each of the measurements in Figures 4 and 5, PNPAm was irradiated with 1450 nm pump wavelength, ∼0.26 mJ pulse energy, 0.38 cm $1/e^2$ radius. Each spectrum exhibits a peak in spectral amplitude at ∼1.5 THz, followed by an absorption dip just above 2 THz. There is significant amplitude up to higher frequencies, however, the response is limited by the pump pulse duration. The fact that the spectra have a similar shape across this broad range of pump wavelengths indicates that the pump group index varies minimally across the same range. A brief description of phase matching relevant for THz generation is discussed in the Supporting Information.

Figure 6b shows the peak-to-peak electric field versus pump wavelength. We see that from 1250 to 1800 nm there is a fairly consistent peak-to-peak THz field strength, with an approximately 10% increase leading to a maximum output at 1565 nm. From 1800 to 2150 nm, the field strength monotonically decreases to roughly half the maximum value. The inset to Figure 6b shows the percent transmission of light through PNPAm.
as a function of wavelength from 500 to 2000 nm. The absorption cutoff at 611 nm is marked with a yellow dot. The dip in the peak-to-peak electric field at 1500 nm can be explained by the absorption of the pump light at 1500 nm.

We next characterized how PNPA generates THz as a function of pump fluence. To extend the range of accessible pump fluences, we used a telescope to reduce the pump spot size to a $1/e^2$ radius of 0.19 cm. With this spot size and a pulse energy of $\sim 0.63 \text{ mJ}$ at 1450 nm, the highest fluence reached was $\sim 5.5 \text{ mJ/cm}^2$. Visual inspection and repeat use of the crystal showed no damage to the crystal. To more carefully characterize the damage threshold, we would need to measure with higher fluences. In Figure 7, we show the pump-fluence dependence of PNPA (690 μm), directly compared to DAST (590 μm) and OH-1 (320 μm) with the same experimental conditions for each crystal. Although there are differences in the thicknesses of the crystals, we picked the highest quality crystals for this direct comparison.

Saturation effects start to appear in PNPA at $\sim 2 \text{ mJ/cm}^2$, as suggested by the dotted red line. We also characterized the generation efficiency of all three crystals, as shown in the inset. These measurements demonstrate that PNPA generates significantly more THz than DAST and OH-1. We see peak-to-peak field strengths approaching 3 MV/cm, as well as a higher efficiency.

Figure 7 shows a clear advantage of using PNPA over DAST and OH-1, however, using a single value to characterize broadband THz generation is inadequate. To further elaborate, Figure 8 shows the THz traces (Figure 8a) and spectra (Figure 8b) of the same crystals recorded at the maximum fluence conditions in Figure 7. PNPA exceeds DAST and OH-1 in both the maximum field strengths achieved and in spectral breadth. DAST and OH-1, respectively, reach peak-to-peak electric field strengths of 2.2 and 1.5 MV/cm, while PNPA reaches 2.9 MV/cm. The combination of an exceedingly high field strength and
large spectral amplitudes across a broad range of frequencies confirms that PNPA should be used as a new standard in high-field THz generation.

We also note that PNPA is more amenable to growing large crystals and polishing smooth surfaces than DAST and OH-1, which can crack more easily during the polishing process. This advantage makes PNPA not only an extremely efficient THz generator, but a more synthetically accessible organic NLO crystal as well.

■ CONCLUSION

In summary, we demonstrate crystallization methods for PNPA that enable access to high quality single crystals for THz generation applications. When comparing PNPA to crystals that have been the standards in high-field THz generation for several years (DAST and OH-1), we find that PNPA outperforms these standards, granting access to much higher electric field strengths (peak-to-peak 2.9 MV/cm for PNPA compared to 2.2 MV/cm for DAST and 1.5 MV/cm for OH-1). PNPA also generates a broad and smooth spectrum from 0 to 5 THz with only one minor absorption at 2 THz, with peak absorption coefficient <200 cm$^{-1}$. We present thickness and wavelength dependent data, showing that PNPA performs well over a broad range of pump wavelengths with minimal changes to the generated spectrum. PNPA also shows the highest THz generation efficiency of an organic NLO crystal to date, exceeding 4%. These results demonstrate the utility of PNPA for THz generation compared to other commercial organic NLO crystals and sets a new standard for high-field THz generation.

■ ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsphotonics.2c01336. Refractive index information, determination of crystal face, table of experimental parameters (PDF)
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