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Organic piezoelectric materials: milestones and potential

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Abstract

Research on the piezoelectric response of biomolecules has intensified following demonstration of open circuit voltages of over 20 V in biopiezoelectric generators. Organic piezoelectric nanotubes, fibers, and micro-islands have been grown and studied; however, the lack of fundamental understanding of the piezoelectric effect in nature hinders the rational design of biomaterials to provide a tailor-made piezoelectric response. Advances in high performance computing have facilitated the use of quantum mechanical calculations to predict the full piezoelectric tensor of biomolecular crystals, including amino acids and small peptides. By identifying directions of high piezoelectric response, the simulations can guide experimental crystal growth, device fabrication and electrical testing, which have led to the demonstration of unprecedented piezoelectric responses in organic crystals on the order of 200 pC/N. These large responses arise from strong supramolecular dipoles, which can be tuned by molecular chemistry and packing, opening new opportunities for the realization of technologically useful piezoelectric devices from renewable materials. The amino acids predicted to exhibit the highest piezoelectric response, such as glycine, hydroxyproline and lysine, are anticipated to be used to engineer highly piezoelectric peptides in the future. With improved scaling of advanced computational methods, such as density functional perturbation theory, the research community can begin to efficiently screen peptide structures for enhanced electromechanical properties. This capability will accelerate the experimental development of devices and provide much-needed insight into the evolution of a hierarchical relation in biological materials starting from strongly piezoelectric building blocks.

Biological materials are low-symmetry, highly ordered structures that lack an inversion center. Thus, linear electromechanical coupling (piezoelectricity) is an inherent functional property of the majority of biomolecules. Bystrov et al¹. describes biological piezoelectricity as being based on complex dipolar properties and dipole–dipole interactions mediated by intricate hydrogen bonding networks with different levels of self-assembly and hierarchy. In the past 60 years, piezoelectricity has been confirmed in a variety of biological materials, such as wood and bone, as well as fibrous proteins such as collagen, chitin, and elastin², which present as highly ordered crystalline molecules in mammalian tissue. Classical piezoelectric principles have been applied to similar uniaxially orientated, bioactive polymers, such as poly (L-lactic acid) (PLLA), poly (γ -benzyl glutamate) (PBG), and cellulose³. These biodegradable polymers have been used as piezoelectric implants to enhance bone formation in both pure and compound forms⁴. Until the 21st century, biological piezoelectricity was not considered technologically significant, with experimental values ranging from 0.1 to 2 pC/N^2 . For comparison, commercially exploited piezoelectrics, such as aluminum nitride (AlN) and lead zirconate titanate (PZT)⁵, show piezoelectric constants of approximately 8 pC/N and 800 pC/N, respectively. A comparison of the piezoelectric strain constants of biological materials to date with a variety of inorganic piezoelectrics is shown in Fig. 1. For further comparisons, readers are directed to the works of Aizen⁶, Shrout⁷, Chorsi⁸, and Wei⁹.

Spurred by demonstration of open circuit voltages of over 20 V in piezoelectric generators made of viruses¹⁰, prawn shells¹¹, and fish bladders¹² (see examples, Fig. 1b),

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biological piezoelectric research has intensified in recent years. Organic piezoelectric nanotubes, fibers and microislands have been grown and studied¹³; however, the lack of fundamental understanding of the piezoelectric effect in biomolecules hinders the design of tailor-made piezoelectric responses in materials engineering. For inorganic piezoelectrics, atomic-scale investigations continue to highlight how a nanoscale understanding of piezoelectricity is necessary to enhance macroscopic electromechanical properties^{14–16}.

Quartz, zinc oxide, and AlN are three classically exploited piezoelectric materials¹⁷. To study and quantify

fundamental piezoelectric properties, researchers have historically focused on single crystals from which the behavior of the bulk material can be understood by the repeated pattern of atomic or molecular-scale unit cells. However, this approach has not been the case for biological piezoelectricity. The majority of experimental investigations into biological piezoelectricity in the past 60 years have been on macroscopic or polymeric samples. This can be attributed in part to the extreme difficulties in crystallizing fibrous and transmembrane proteins, but a large number of amino acids and peptides have been crystallized since the advent of X-ray diffraction¹⁸ and, in



particular, studied for their nonlinear optical (NLO) properties¹⁹. Lysozyme, a well-known globular protein found in tears and saliva, was only recently reported to exhibit piezoelectric properties in crystalline forms²⁰. In addition to lysozyme, one of the nucleobases, thymine, was shown to be ferroelectric and hence piezoelectric in the crystalline form²¹. Biological materials present new challenges in the field of piezoelectricity, most notably predicting the role of water in modulating the measured dielectric constants and thus piezoelectric properties. A further bottleneck is the inherent softness of biological materials in a field where measurements have traditionally required application of external mechanical forces. The small size of the majority of biomolecular crystals also limits how electrical contact can be made-particularly compared to large inorganic piezoceramics that can be sliced, polished and electroded with relative ease.

Single crystal studies present a number of advantages for investigating piezoelectricity in any material, including biological structures. First, in terms of theoretical methodology, quantum mechanical modeling most often begins with the unit cell. Density functional theory (DFT)²² utilizes periodic boundary conditions to simulate bulk material behavior and can quantify material physical properties of crystals, including the dielectric, elastic, and piezoelectric constants. By studying biomolecular crystals in this way, the predicted physical properties can be directly related to single crystal experiments, allowing effective screening of organic crystals for experimental investigation. A combination of modeling and characterization can provide much-needed insight into how piezoelectric properties are modulated by unit cell properties, such as dipole moments, molecular packing, and composition (Fig. 1a). Figure 1a shows that crystallizing biomolecules creates a network of unit cell dipoles identical to the mechanisms of classical inorganic piezoelectrics, which allows for biological single crystals to easily fulfill the role of piezoceramics, e.g., in stack actuation. Single crystals can often be grown quite easily from aqueous solutions, and the quality and shape of organic single crystals can be modulated using additives or pH buffers. This simple idea that the properties of single crystals can be modified chemically to enhance their properties suggests that this could be extended to piezoelectric properties using theoretical predictions to guide experiments.

Lemanov first suggested in 2000^{23} that the key to understanding biological piezoelectricity was to study the 20 protein amino acids that are used to build all biological materials. Amino acids crystallize primarily in lowsymmetry orthorhombic and monoclinic space groups (Fig. 2). These space groups are noncentrosymmetric, which lend themselves to piezoelectric and NLO properties, with the exception of the largest amino acid, Ltryptophan, and glycine. L-tryptophan crystallizes with the lowest *P*1 crystal symmetry²⁴. Glycine is not chiral thus it does not have an enantiomorphic mirror image, and it crystallizes in three distinct crystal symmetries: α , β , and γ^{25} . The only amino acid structure that precludes piezoelectricity is α -glycine. The highest responses



measured so far in amino acid single crystals are 178 pC/ N and 25 pC/N in β -glycine and hydroxy-L-proline, respectively^{26,27}.

In 2012, Lemanov²⁸ fired high frequency electrical pulses at crystalline amino acid powders (not single crystals) and found that DL-alanine and γ -glycine had the strongest piezoelectric response of the amino acids studied. This result is consistent with recent quantitative investigations into longitudinal piezoelectricity in these crystals,^{26,29} which determined that both DL-alanine and *y*-glycine crystallize in unique space groups that allow for a longitudinal d_{33} constant of 10 pC/N. Lemanov did not detect piezoelectricity in L-cysteine or L-isoleucine powders, which are noncentrosymmetric in their single crystal form. This result highlights the importance of using crystallized amino acids as opposed to powders when studying chiral molecules. A false piezoelectric signal was also measured in α -glycine, which was attributed to contamination from other noncentrosymmetric polymorphs. Furthermore, an orientation dependence of the piezoelectric response of glycine powder was noted, which has since been explained quantitatively 26 . When γ -glycine crystals are rotated sequentially by 90° between electrodes (so that their crystallographic a, b, and c axes become perpendicular to the plane of electrical contacts), their measured piezoelectric constant changes from approximately 1 to 2, and finally to 10 pC/N^{26} . Prior to Lemanov's work in 2000, the only experimental investigation into amino acid piezoelectricity had been performed by Vasilescu et al. in 1970.³⁰ The noncentrosymmetry of the amino acids was confirmed in 1967 by NLO methods³¹. Amino acid compound crystals have been more widely studied for their piezoelectric properties, including L-arginine phosphate (LAP)³² and triglycine sulfate $(TGS)^{33}$. Doping of centrosymmetric α -glycine crystals with other L-amino acids also leads to crystals with piezoelectric properties³⁴. The peptide that has attracted the most attention for its piezoelectric properties is di-phenylalanine (FF), which is easily influenced by external electric fields in aqueous solution³⁵. FF peptide nanotubes have been confirmed to be piezoelectric and pyroelectric^{1,36,37}. These peptides also demonstrate high stability and a large Young's Modulus, which has made them potential alternatives to a number of piezoelectrics widely used in consumer electronics. Recent work has shown that FF nanogenerators can generate voltages of 0.6-2.8 V, which places them alongside glycine and alanine as ideal candidates for self-powered brain implants and pacemakers³⁸.

New challenges arise for predictive modeling as we continue up the biomolecular hierarchy, specifically to peptide and protein crystals (Fig. 3). Although trends in piezoelectric charge tensors can be seen in the sequences of small peptide crystals²⁷, it is unrealistic to track, for example, elastic constants through the hierarchy of biomolecular crystals. The amino acids predicted to exhibit the highest piezoelectric response, such as glycine and lysine, are thus anticipated to be used to engineer highly piezoelectric peptides in the future. With improved scaling of advanced quantum mechanical methods, such as density functional perturbation theory (DFPT, peptide structures can be efficiently screened for enhanced electromechanical properties.

To conclude, piezoelectricity is a ubiquitous yet underexploited property of the majority of biomolecules. Studying biomolecules in their crystallized form allows for the understanding of their electromechanical behavior in terms of unit cell properties. A single crystal approach is needed to fully understand and exploit biological piezoelectricity, allowing for high-throughput, simulationguided experiments. A solid-state physics perspective on biomaterials will aid the adaptation of biological crystals in high performance sensors, actuators, and energy harvesters. Biological piezoelectricity also holds unique potential for in vivo sensing and drug delivery, which is diversifying current research beyond the replacement of lead-based piezo materials and requires bold advances in predictive modeling of the electronic structures of large biomolecular crystals and supramolecular assemblies.

Conflict of interest

The authors declare that they have no conflict of interest.

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