

Electric-field-stimulated protein mechanics

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The internal mechanics of proteins—the coordinated motions of amino acids and the pattern of forces constraining these motions—connects protein structure to function. Here we describe a new method combining the application of strong electric field pulses to protein crystals with time-resolved X-ray crystallography to observe conformational changes in spatial and temporal detail. Using a human PDZ domain (LNX2^{PDZ2}) as a model system, we show that protein crystals tolerate electric field pulses strong enough to drive concerted motions on the sub-microsecond timescale. The induced motions are subtle, involve diverse physical mechanisms, and occur throughout the protein structure. The global pattern of electric-field-induced motions is consistent with both local and allosteric conformational changes naturally induced by ligand binding, including at conserved functional sites in the PDZ domain family. This work lays the foundation for comprehensive experimental study of the mechanical basis of protein function.

The fundamental biological properties of proteins—binding, catalysis and allosteric communication—emerge from a global pattern of interactions between all constituent atoms. Often, this pattern is organized in the tertiary structure so as to produce the concerted motions of amino acid residues, defining transitions between a small number of functional states. This conformational cycling within proteins and protein complexes draws analogies to macroscopic machines¹ and lies at the heart of many biological processes: DNA replication², metabolism^{3,4}, transport⁵, cellular motility⁶ and signal transduction⁷. Even without conformational changes, functional states of proteins can have a different pattern and extent of rigidity^{8,9}—entropic variations that also influence state transitions¹⁰. Thus, the biology of proteins is deeply connected to their mechanics: the motions a protein can perform and the forces constraining these motions. As in macroscopic machines¹, a comprehensive description of internal mechanics is the key to explaining how structure leads to function¹¹. Unlike conventional machines, however, proteins are marginally stable evolved materials whose mechanics are governed by weak, heterogeneously cooperative interactions for which we as yet have no good physical models.

Current biophysical methods provide an incomplete basis for making mechanical models of proteins. NMR spectroscopy¹² and room-temperature crystallography¹³ provide information on the structure and dynamics of local environments of atoms, and have been used to characterize weakly populated excited states of proteins^{3,13,14}. However, the complexity of disentangling conformational transitions occurring on multiple timescales, the difficulty of directly seeing collective motions, and the inability to generally relate the measured parameters to physical forces limit our understanding. Single molecule force spectroscopy can relate global conformational transitions to applied forces, but is limited in providing the atomic detail required to define the underlying intramolecular mechanics¹⁵. Time-resolved crystallography (TRX) offers, in principle, a direct route to observing concerted motions with high temporal and spatial resolution¹⁶. However, TRX traditionally relies on photoexcitation of bound chromophores to induce motions in proteins¹⁷. Such excitation is not generally applicable, acts at a fixed location, is not tunable, and deposits an amount of

energy that far exceeds the typical energetic changes involved in protein conformational transitions.

Here, we describe a new method for studying protein mechanics and its application to a model system—a PDZ domain—which shows both local and allosteric functional properties¹⁸. The method, electric field-stimulated X-ray crystallography (EF-X), combines the use of strong electric field pulses to drive motions within protein crystals with simultaneous readout by fast X-ray pulses. EF-X satisfies the key characteristics required for a general mechanical analysis of proteins: (1) the application of forces of controlled magnitude, direction and duration; (2) the existence of defined, well-distributed actuators (the charges) on which the forces act; and (3) readout of conformational changes with high spatial and temporal resolution. We show that EF-X can reveal protein motions associated with biological function and permits direct refinement of the atomic structures of low-lying excited states. This work initiates a path towards a full description of protein mechanics.

Theoretical and practical considerations

The idea of EF-X is simple; many elementary charges and local dipoles are present in proteins (Fig. 1a), and with the application of sufficiently large external electric fields, it should be possible to exert forces on them that cause motions of atoms throughout the protein structure. If the electric field can be applied in conjunction with timed X-ray diffraction in protein crystals, it should be possible to observe all of these motions in high spatial and temporal detail (Fig. 1b). To implement the idea, we began with a few design considerations. Theoretical calculations suggest that electric field strengths of $\sim 1,000,000 \text{ V cm}^{-1}$ are in the right range to drive subtle motions of atoms within proteins that can be observed through high-resolution diffraction methods (Methods and Extended Data Table 1). Fields of 1 MV cm^{-1} are dangerously large from a laboratory point of view, but are close to physiological; for example, 0.125 MV cm^{-1} corresponds to $\sim 100 \text{ mV}$ across a cell membrane. Such voltages influence conformational transitions in proteins such as ion channels¹⁹ and G-protein-coupled receptors²⁰, and are consistent with biological relevance. In general, the basic premise of EF-X is that features corresponding to the biologically relevant reaction coordinate(s) of proteins are enriched in the low-lying energetic states around

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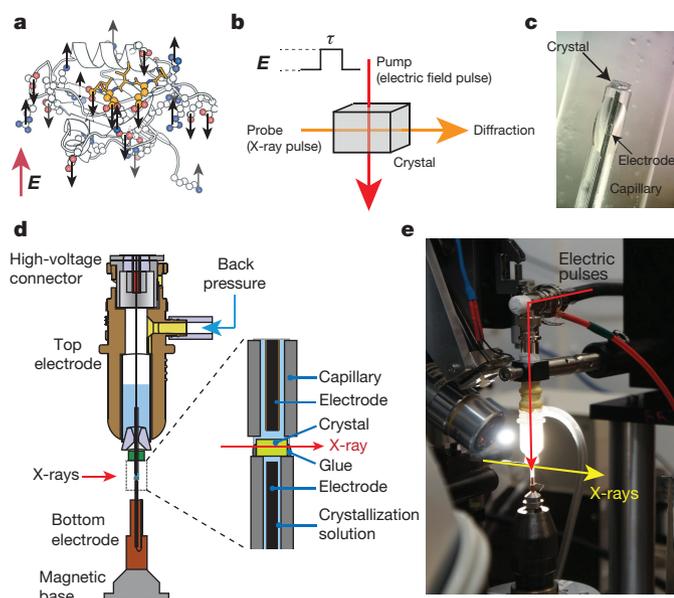


Figure 1 | EF-X principles and implementation. **a**, A sampling of charged residues (in CPK colours) in LNX2^{PDZ2} (Protein Data Bank (PDB) accession 2VWR), exemplifying potential actuators for applied electric fields (E , in red). Ligand is in yellow. **b**, EF-X involves stimulation of motions in protein crystals by an applied electric field (E) of duration τ (the ‘pump’), and readout by much faster X-ray pulses (the ‘probe’). **c**, An LNX2^{PDZ2} crystal mounted across the orifice of a glass capillary, filled with crystallization solution and a metal electrode. The crystal is sealed onto the capillary by an electrically insulating glue. **d**, The crystal is mounted on the bottom electrode and the high voltage is delivered from a top electrode through a liquid junction composed of crystallization solution. Controlled back pressure on a reservoir of solution in the top electrode keeps the crystal continuously hydrated. **e**, A view of the assembled experimental apparatus.

the ground state, and that forces imposed by $\sim 1 \text{ MV cm}^{-1}$ electric fields represent an effective strategy to bias and expose these states.

Practically, there are several experimental complications (Supplementary Information IA). Of these, the main one is crystal heating caused by electric-field-induced flow of ionic currents through solvent channels. If sufficiently large, this effect leads to dielectric breakdown, arcing, destruction of the crystal, and a dramatic end to the experiment (Supplementary Video 1). However, calculations with estimated conductivities of protein crystals²¹ and typical crystallization solutions suggested that electric fields on the order of 1 MV cm^{-1} should be tolerated for pulse durations up to microseconds (Supplementary Information IA). Together with rise-time limits of our current high-voltage system ($\sim 10 \text{ ns}$) and electrode design, this defines a window of timescales for these experiments at present (Extended Data Fig. 1a). These limits can be extended through further technical development.

On the basis of these considerations, we built a custom setup for room-temperature X-ray diffraction of protein crystals under strong electric field pulses on the sub-microsecond timescale (Fig. 1c–e, Methods and Extended Data Fig. 1). Protein crystals are sandwiched between two glass capillaries filled with crystallization solution and containing metal wires that serve as electrodes (Fig. 1c). The crystal is fixed by an electrically insulating glue to the bottom (ground) electrode and the high-voltage pulse is introduced from a top electrode through a liquid contact with the crystal (Fig. 1d and Supplementary Video 2). See Methods and Extended Data Fig. 1 for design details. This electrode system was integrated into a synchrotron X-ray facility designed for time-resolved crystallography (BioCARS²², Advanced Photon Source; Fig. 1e).

Application of EF-X to the PDZ domain

As an initial model system, we chose the second PDZ domain of the human E3 ubiquitin ligase LNX2 (LNX2^{PDZ2})²³ (Fig. 2a). PDZ domains

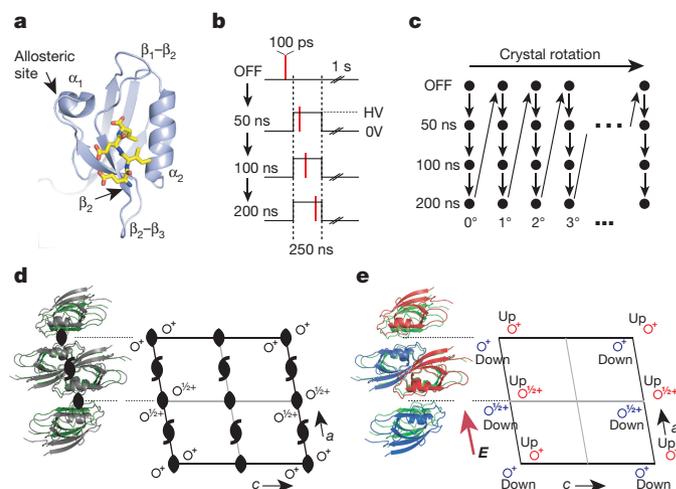


Figure 2 | An EF-X experiment in the LNX2^{PDZ2} domain. **a**, LNX2^{PDZ2} binds target ligands (in yellow) in a groove between the β_2 and α_2 segments. The binding site is coupled to allosteric sites on the β_2 – β_3 segment and the α_1 – β_4 segment (through the β_1 – β_2 loop and α_1 helix). **b**, Data collection involves four sequential X-ray exposures for each crystal orientation: no voltage (OFF), and three time delays (50, 100, 200 ns) after onset of the voltage pulse. One second is allowed between pulses for crystal cooling. HV, high voltage. **c**, The protocol in **b** is repeated for a series of crystal rotations to collect a full diffraction data set. **d**, LNX2^{PDZ2} crystallizes in the C2 space group, which includes two kinds of rotational symmetry (black symbols); this results in four molecules per unit cell and one molecule per asymmetric unit. **e**, With the electric field E (applied along the a dimension), all rotational symmetry is broken. This results in a new unit cell with two molecules per asymmetric unit (red and blue)—one experiencing $+E$, and one experiencing $-E$.

are 90–100-residue proteins that generally bind the C termini of target proteins between the α_2 helix and β_2 strand²⁴. Previous data demonstrate the existence and functional relevance of allosteric coupling of the ligand-binding site to a few distant surfaces²⁵, especially the α_1 helix^{26,27} and the β_2 – β_3 loop²⁸ (see Supplementary Table 1). Otherwise, LNX2^{PDZ2} is a typical protein, with no special features that compromise the generality of this study. Specifically, LNX2^{PDZ2} has no known functional voltage dependence, providing a test that EF-X can be generically used in the context of randomly available formal and partial charges for analysis of protein mechanics.

We performed EF-X experiments on LNX2^{PDZ2} with voltage pulses of 5–8 kV to 50–100- μm -thick crystals, resulting in field strengths of ~ 0.5 – 1 MV cm^{-1} . The pulse durations ranged from 50 to 500 ns, and diffraction was collected with single 100 ps X-ray pulses. The pulse protocol permits us to examine the atomic structure before the electric pulse (voltage-OFF data set) and at any specified time delay after initiation of the electric pulse (voltage-ON data set) (Fig. 2b, c). The OFF data set provides a reference structure for study of electric-field-induced effects. As predicted by our calculations, LNX2^{PDZ2} crystals (and other protein crystals) readily tolerated hundreds of 100–500 ns electric field pulses of $\sim 1 \text{ MV cm}^{-1}$ and X-ray pulses without substantial loss of diffraction (Supplementary Table 2 and Extended Data Fig. 2). We collected a time series from a single LNX2^{PDZ2} crystal, consisting of an OFF data set and ON data sets at 6 kV and at 50, 100 and 200 ns delays from the rising edge of the electric field pulse (Fig. 2b, c and Supplementary Table 3); the variability in timing is less than 1 ns and is therefore negligible given the timescale of this experiment.

Breaking symmetry

An important analytic tool comes from understanding how the electric field affects the symmetry S of the crystal lattice. In general, the unit cell of a protein crystal can be constructed from a set of symmetry operations $\{S\}$ —combinations of translations and rotations—that define its

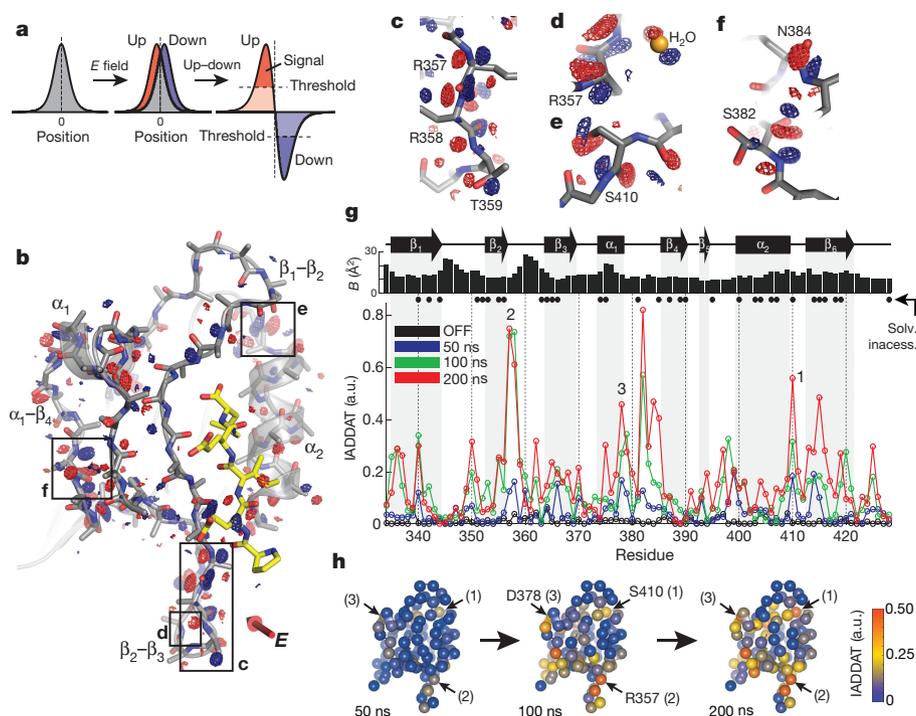


Figure 3 | The up-down internal difference analysis. **a**, In the simplest case, the electric field will shift the electron density distribution for an atom in the up and down molecules (red and blue, respectively) in opposite directions around its centroid in the voltage-OFF molecule (grey) (left and middle). Subtracting the up and down densities and applying a noise threshold (right), we expect peaks of positive (red) and negative (blue) difference density surrounding an atom in the OFF state—the hallmark of an electric-field-induced motion. **b**, The up-down internal difference map for a ‘front’ view of LNX2^{PDZ2}, with regions highlighted in c–f boxed. The red three-dimensional arrow indicates the direction of the electric field, and bound ligand is in yellow. Maps are contoured at +3.5 (red) and –3.5 (blue) σ_{OFF} and, for clarity, are displayed within

characteristic space group. For example, the LNX2^{PDZ2} crystals have space group $C2$, which, in addition to translational symmetry, has two kinds of rotational symmetry elements (Fig. 2d). As a consequence, there are four symmetric LNX2^{PDZ2} monomers per unit cell. What happens if an electric field is applied in a certain direction? Clearly, protein molecules in the crystal lattice with different orientations relative to the field will undergo different changes and will no longer be symmetric. The general rule is that any crystal symmetry operator S that does not preserve the orientation of the electric field E will be violated (‘broken’: $S \cdot E \neq E$). For the LNX2^{PDZ2} experiment, the electric field breaks all the $C2$ rotation symmetry operators (Fig. 2e). Now, the four LNX2^{PDZ2} molecules in the unit cell are no longer equivalent, and symmetry is reduced such that two molecules see the field in one direction (we will refer to these molecules as ‘up’), and two see the field in the opposite direction (the ‘down’ molecules). In essence, if the up molecule experiences +6kV, the down molecule experiences –6kV, and so the force acting on otherwise equivalent atoms in these structures is opposite in direction. Although it need not be strictly symmetric, we would naively expect this to cause an opposite motion of atoms from their mean positions in the OFF state (Fig. 3a).

This breaking of symmetry provides a powerful way to study the effect of the electric field on the protein structure. We can compare the up and down molecules within the unit cell, an internally controlled experiment that isolates the effect of the electric field on atoms. In contrast, artefacts due to radiation damage and heating are insensitive to the direction of the electric field and cancel out in this analysis (see Methods). In crystallographic terms, we compute an internal difference Fourier map in which we subtract the up and down electron densities

a 1.8 Å shell around main chain + C β atoms (see PyMOL session S1 for full map). **c–f**, Examples of electric-field-induced motions—opposing red and blue density—for main-chain, side-chain and solvent atoms. **g**, The IADDAT for the up-down molecules as a function of LNX2^{PDZ2} primary structure and time (blue, green and red traces). The OFF difference density (black) indicates the noise in the analysis. The graphs above indicate buried residues (solvent accessibility <0.15 (Solv. inaccess.)) and refined isotropic B -factor for the voltage-OFF model. a.u., arbitrary units. **h**, The time evolution of the electric-field-induced effects mapped on the tertiary structure of LNX2^{PDZ2}. Spheres indicate C α positions and colours IADDAT.

(Fig. 3a). In such a map, the hallmark of an electric-field-induced structural effect is to see peaks of opposite sign around the position of an atom in the voltage-OFF state (red and blue, Fig. 3a).

The up-down map shows pervasive evidence of electric-field-induced atomic motions (Fig. 3b–f). Just as proposed, we observe shifts of backbone, side-chain and solvent atoms in opposite directions between the up and down molecules (Fig. 3c–f). The structural response is distributed broadly over the protein tertiary structure, in both core and surface sites, with some of the strongest signals around the β_2 – β_3 , α_1 – β_4 and α_2 – β_6 segments (Fig. 3b). To examine the response quantitatively, we integrated the absolute difference electron density above a noise threshold (IADDAT) in a volume shell around the protein backbone²⁹ (Fig. 3g). The up-down effect in the OFF state provides a measure of noise (black trace, Fig. 3g). By comparison, we observe a robust signal in the presence of the electric field that evolves over time from 50 ns to 200 ns (Fig. 3g, blue, green and red traces, and Fig. 3h). The electric-field-induced motions do not simply reflect solvent exposure or thermal (B) factors related to positional disorder (for all cases, $P > 0.1$, Fisher Z -test; see Methods) (Fig. 3g). Many of the affected residues do not have formally charged side chains, indicating that they move due to local dipoles or due to structural coupling with other charged residues. An extensive statistical validation of signal to noise is presented in Extended Data Figs 3a–c and 4, Supplementary Tables 4–6 and Supplementary Information IB.

The signal evolves heterogeneously over the structure (Fig. 3g, h), with some regions moving over the full time period (for example, peaks 1 and 3), and others complete at intermediate times (for example, peak 2). This variation in characteristic timescales of motion in different

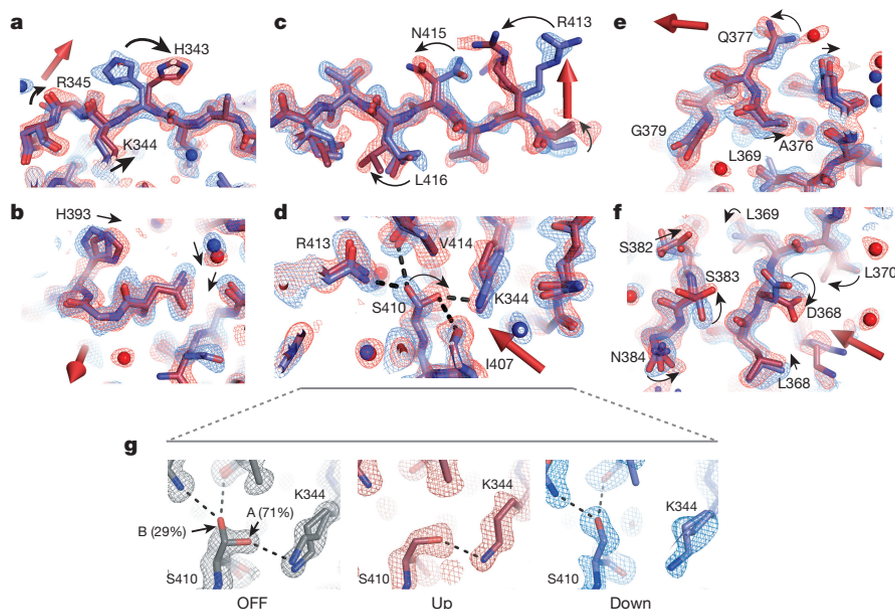


Figure 4 | A gallery of electric field-induced structural effects.

a–f, Refined models and associated $2F_o - F_c$ electron density contoured at 1.5σ for the up (red) and down (blue) LNX2^{PDZ2} structures (6 kV, 200 ns delay); the direction of the electric field is indicated by the three-dimensional arrow. The data show examples of rotamer flips (H343; **a**), continuous displacements (H393; **b**), potentially coupled rotamer flips (R413, N415, L416; **c**), rearrangements of hydrogen bonding (S410; **d**), motions of secondary structure elements (the α_1 helix; **e**), and complex combinations of these effects (**f**). Per sign convention, atoms coupled to a positive charge

in the up model would move in the direction of the field and in the down model, against the field. Motions occur at solvent-exposed (**a–c**, **e**) and -buried (**d**, **f**) regions (PyMOL sessions S2–S4 and Extended Data Fig. 6). **g**, S410 shows partial occupancy in two rotameric states (marked A and B) in a 1.1 Å room-temperature ground-state structure (OFF, Extended Data Table 3). These states are biased by the electric field such that the up and down models each adopt one of the two ground-state configurations (middle and right). Maps are contoured at 1.5σ . See Extended Data Fig. 7 for more examples.

regions of the structure is a property that, with further study, could be deeply informative about the underlying pattern of forces between amino acid residues. A broad analysis of crystal growth conditions, diffraction quality and symmetry suggests that many protein crystals should be amenable to the EF-X experiment, including use of the up–down difference method (Supplementary Information ID).

Modelling of excited states

We refined atomic structures of the up and down states of the LNX2^{PDZ2} domain at 200 ns from the onset of the electric field. Since the field only subtly biases the ground state conformation, we carried out refinement against extrapolated structure factors (ESFs)^{30,31} (Extended Data Table 2). The ground state (OFF model, Extended Data Table 3) was used as a starting point, with progress supported by R factors ($\Delta R_{\text{work}} = -6.96\%$, $\Delta R_{\text{free}} = -5.92\%$; Methods, Extended Data Fig. 5 and Supplementary Information IB). Propagation of errors suggests that the ESF structures at 200 ns have an effective resolution of 2.3 Å.

The structures demonstrate electric-field-induced perturbations of nearly every type of physical interaction throughout the protein structure—induction of side-chain rotamer flips (Fig. 4a), continuous displacement of backbone atoms, side chains and bound waters (Fig. 4b), propagated rotamer shifts suggesting collective motions through the structure (Fig. 4c), breaking and re-forming of hydrogen bonds (Fig. 4d), global motions of entire secondary structure elements (Fig. 4e), and complex coordinated changes in large regions (Fig. 4f). Extended Data Fig. 6 shows additional examples. For some residues, the electric field biases the occupancy of pre-existing alternate conformational states in the voltage-OFF structure (Fig. 4g and Extended Data Fig. 7). These residues are differentially forced into either of the alternative configurations depending on the direction of the applied field. Thus, rather than inducing non-physiological states, EF-X appears to expose low-lying conformational states that are energetically near to the ground state.

These data validate the broad goals of EF-X—to globally perturb and record subtle motions in a mechanistically unbiased manner at

atomic resolution. A key feature is the ability to actively populate and directly model the structures of low-lying excited states around the ground states of protein molecules, the configurations most likely to be relevant over the functional reaction coordinate. In addition, the ability to collect data sets at various time delays after the initiation of the electric field pulse means that we can observe these motions as they happen in time and make experimental movies of the temporal evolution of protein motions¹⁷.

The biological relevance of stimulated motions

We asked what the electric-field-induced motions tell us about the biology of the PDZ domain. The backbone motions accumulate in four parts of the protein—the α_1 helix and the β_1 – β_2 , β_2 – β_3 and α_2 – β_6 segments—all known to be functionally coupled to ligand binding (Fig. 5a, b). The partially buried α_1 helix and the α_1 – β_4 surface form the central components of allosteric communication in PDZ domains^{27,32–35}, and residues in these regions undergo systematic electric-field-induced shifts and rotameric transitions (Figs 4f and 5a). In addition, the β_1 – β_2 and β_2 – β_3 segments move and become more ordered (Extended Data Fig. 6d), transitions reminiscent of ligand-induced changes in many PDZ domains^{24,25}. Finally, the electric-field-induced switching of S410 between two hydrogen-bonding networks (α_2 – β_6 region, Fig. 5d, g) positions a conserved buried cationic residue (K344) in the ligand-bound configuration in several PDZ homologues³⁶ (Extended Data Fig. 6e).

To test the relationship of electric-field-induced motions to PDZ function rigorously, we analysed the ligand-induced displacements of main-chain atoms averaged over 11 diverse homologues of the PDZ family (Fig. 5c, e). Ligand-induced motions shared by these homologues are most pronounced in the β_1 – β_2 , β_2 – β_3 and α_2 – β_6 segments, and in the α_1 and α_2 helices (Fig. 5c), comprising most regions with electric-field-induced motions. These regions are also linked by the protein sector^{26,37}—a group of amino acid positions that statistically co-evolves in the entire PDZ family—suggesting that the pattern of ligand-induced motions is an evolutionarily conserved feature in

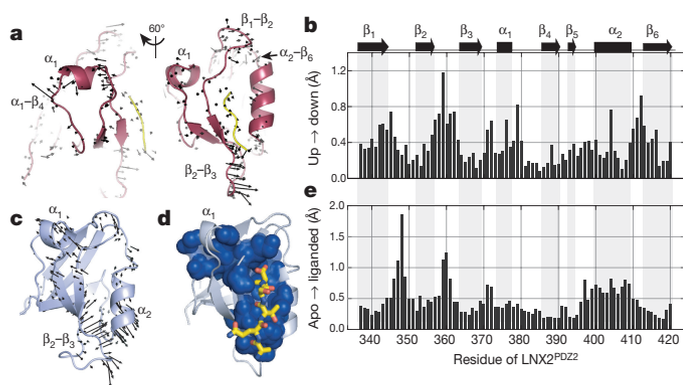


Figure 5 | The relationship between electric-field-induced conformational change and PDZ function. **a**, Two views of the electric-field-induced structural changes in LNX2^{PDZ2} (6 kV, 200 ns data set), with vectors representing the displacements of main-chain atoms transitioning from the up to down models (enlarged $\times 5$ for clarity). The motions are most prominent in the β_1 – β_2 , β_2 – β_3 , α_1 and α_2 – β_6 regions. **b**, The mean displacements of backbone atoms per residue between the up and down states. **c**, Conserved motions of backbone atoms due to ligand binding (apo to liganded) in high-resolution structures of 11 diverse homologues of the PDZ family (vectors enlarged $\times 10$). The motions occur in similar regions as in **a**, but also include the α_2 helix. **d**, The protein sector (blue spheres), a group of coevolving amino acid positions in the PDZ protein family (PFAM 27.0 (ref. 39)); the sector connects the ligand-binding pocket to the β_2 – β_3 segment and to the α_1 – β_4 surface through the β_1 – β_2 loop and the α_1 helix. **e**, The median ligand-induced displacements of backbone atoms per residue (LNX2^{PDZ2} numbering) in the ensemble of 11 PDZ homologues. Statistical comparison with that for the up to down transition (**b**) shows a significant correlation ($P < 0.001$, Fisher Z-test).

the PDZ domain (Fig. 5d). Overall, the pattern of conserved apo to liganded displacements (Fig. 5e) shows a highly significant correlation ($P < 0.001$, Fisher Z-test) with the electric-field-induced up to down motions (Fig. 5b). This result is particularly meaningful because, in principle, ligand binding and electric fields could impose forces in a protein structure in a manner completely distinct from each other, and the comparison reflects an experiment at just one field strength, orientation, and time delay. Thus, EF-X samples motions in the protein structure that are enriched in its biologically relevant mechanical modes.

From structure to mechanics

A central missing tool in our study of proteins is a method to stimulate and record biologically relevant motions over a broad range of time-scales and with atomic resolution. We show that strong but physiological electric fields can be used to examine a wide range of functional conformational changes within a protein. With further development, we expect that EF-X can be broadly used to investigate the structural basis of protein function (see Supplementary Information ID, IE). It will be of interest to extend EF-X to broader timescales of motions (a matter of further engineering, Extended Data Fig. 1a), to characterize motions in proteins with complex multistate conformational changes, and to study the structures of membrane proteins under physiological electric fields.

However, to go beyond the descriptive level of motions to the underlying physics, it is necessary to infer the spatial distribution of forces, and energies, associated with the observed conformational transitions. In this regard, it is informative to compare EF-X with single-molecule force spectroscopy¹⁵. An electric field of 1 MV cm^{-1} (or 10^8 N C^{-1}) exerts 16 pN per elementary charge, a force sufficient to unzip a leucine zipper protein³⁸. Thus, an exciting prospect is to obtain direct force and free energy estimates for both gradual and discrete conformational changes as in force spectroscopy, but with the atomistic detail and temporal resolution made possible by EF-X. This goal is complicated by the cooperative action of amino acids, but EF-X provides a potential

path to address this problem as well. We can collect EF-X data while varying the duration, orientation, spatial pattern and magnitude of applied forces and statistically group residues that move together into collective modes. These modes may represent the basic mechanical units underlying protein function.

This initial report of EF-X does not yet present a simple, turnkey method. Crystal handling, electrode design, data analysis and structure refinement all leave substantial room for improvement. In addition, the analysis of effects induced by one field orientation and at one time-scale is just a starting point for a full description of relevant motions. However, this work provides an experimental foundation for building good physical models for proteins, the critical link between structure and function.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Supplementary Information is available in the online version of the paper.

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Author Information Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to R.R. (rama.ranganathan@utsouthwestern.edu).

METHODS

System design and safety. The design of the experimental system is based on simple physical considerations. An applied electric field E will impose a force on net charge q to cause a displacement Δx along the field. For any residue (or other group of atoms), we can associate a transition dipole moment $\Delta\mu = \sum_i q_i \Delta x_i$, with each motion, where i is an index over atoms (in units of elementary charge times distance (eÅ); $1 \text{ eÅ} \approx 4.8 \text{ D}$). The energetic effect due to the electric field is $-\Delta\mu \cdot \Delta E$, and its significance depends on how it compares to thermal energy $k_B T$; for example, a weakly populated excited state increases in occupancy by ~ 2.7 fold when its energy relative to the ground state is lowered by $1 k_B T$. As shown in Extended Data Table 1, fields of $\sim 1 \text{ MV/cm}$ are in the right range for our purpose. On the basis of this, we designed a $\pm 10 \text{ kV}$ power supply (Spellman HV) which charges a pulse generator (IXYS Colorado), and from which high-voltage (HV) pulses are triggered by a TTL signal from the synchrotron signal processing hardware. This establishes precise timing between X-ray and electric field (EF) pulses. Integrity of the conductive path to the tip of the capillary and its associated propagation delay were determined using an HV probe.

We designed a number of safety features. Custom RG-11 high-voltage cables (Gater Industries) were high-potential tested by the power group at the Advanced Photon Source and were approved for use up to 8 kVDC . EF pulses were generated in 'half-bridge' mode, where residual charge stored by the HV cables after an EF pulse is drained through a large capacitor connected to ground. The counter electrode was designed to avoid any path through air of less than 1 cm to the grounded cable connector exterior. The inhibit feature of the power supply was connected to an interlock system at the beamline facility, ensuring that the system is de-energized upon personnel entry into the beamline hutch. Power supply voltage and the counter electrode backpressure were controlled remotely using a network-connected microcontroller and custom software.

Electrode construction. The RG-11 cable is terminated on one side with an HV connector (LEMO) (for the pulse generator) and on the other side with an SHV connector (for the housing of the top counter electrode) (Extended Data Fig. 1). The housing was prototyped in-house using a three-dimensional printer (Formlabs, Somerville, MA) and custom fabricated commercially (PolyJet technology, PartSnap, Irving, TX). The housing contains a cylindrical glass insert fitted with a silicone gasket and a thin metal wire ($75 \mu\text{m}$ diameter, Cooner Wire, Chatsworth, CA) with a dielectric coating, except at the tip. The wire was guided to the crystal through a glass capillary (0.5 or 1.0λ (140 or $200 \mu\text{m}$, respectively) orifice, Drummond Scientific, Broomall, PA). The electrode housing was filled with crystallization solution and contains a small port (blue arrow, Extended Data Fig. 1b) that allowed for computer-controlled backpressure for slow infusion of liquid through the top electrode to maintain crystal hydration. Bottom electrodes were prepared from glass capillaries (0.25λ , Drummond Scientific) with a $\sim 100 \mu\text{m}$ orifice, cut in half and aminosilanized at the tip surface to improve adhesive capacity. A $75 \mu\text{m}$ diameter uncoated stainless steel wire (Cooner Wire) was threaded until just below this orifice. The capillary was inserted in a reusable goniometer base (MiTeGen, Ithaca, NY) and soaked, in inverted position, in crystallization solution.

Protein expression, purification and crystallization. For LNX2^{PDZ2}, we obtained an expression strain (BL21(DE3)-R3-pRARE2) and plasmid construct (pNIC28-LNX2^{PDZ2}) from the Structural Genomics Consortium (SGC)⁴⁰ (<http://www.thesgc.org>; construct identifier LNX2A-c033). pNIC28-LNX2^{PDZ2} includes residues 336–424 from *Homo sapiens* LNX2, with the F338L mutation described by the SGC, an N-terminal cloning artefact (334–335), and a C-terminal ligand motif Glu-Ile-Glu-Leu (425–428). LNX2^{PDZ2} protein was expressed as an N-terminal hexahistidine fusion in BL21(DE3)-R3-pRARE2 and purified by nickel affinity chromatography (Ni-NTA agarose, Qiagen), cleavage of the TEV tag by 1 U ProTEV per $50 \mu\text{g}$ protein during dialysis into 50 mM HEPES pH 7.5, 500 mM NaCl, 5% glycerol, 0.5 mM TCEP, size exclusion chromatography, and concentrated to 20 mg/ml for storage. Two protocols yielded suitable crystals. In the first, 3.5 mg/ml protein was dialysed twice (12 and 6 h) against 3 l 5% glycerol, and crystallized by the hanging drop vapour diffusion method in 19% PEG-300, 48 mM citric acid, 35 mM NaH_2PO_4 and 5% glycerol at 20°C . Drops were set up by mixing $0.55 \mu\text{l}$ protein and $1.0 \mu\text{l}$ buffer. In the second protocol, concentrated protein was diluted to 3.5 mg/ml with 10% glycerol and crystallized by hanging drop vapour diffusion in 27 – 31% PEG-300, 43 mM citric acid and 35 mM NaH_2PO_4 at 20°C (drops, $1.0 \mu\text{l}$ protein and $1.0 \mu\text{l}$ well solution).

Crystal mounting. Crystals were manually mounted under a stereomicroscope across the orifice of the pre-soaked bottom electrode, attached to a magnetic goniometer base. Sylgard 184 (Dow-Corning) was prepared to just before full curing and applied around the crystal using a piece of monofilament fishing line (Cajun Line, $0.012''$ diameter, Zebco, Tulsa, OK), taking care to not overcoat the crystal. A MiTeGen polyester sleeve containing $15 \mu\text{l}$ of $50/50$ crystallization solution and water at one end, was slid over the electrode to maintain suitable vapour pressure

for the crystal. The mounted electrode system was placed on the goniometer and the final experimental configuration (Extended Data Fig. 1e, g), was achieved in three steps: (1) coarse relative positioning of the two electrode system using an XYZ translation stage (Thorlabs), (2) cutting the MiTeGen sleeve to expose the crystal, and (3) rapid, camera-guided approach of the top counter electrode until a liquid junction with the crystal was established (Supplementary Video 2).

Data collection and reduction. EF-X data were collected at BioCARS (14-ID) at the Advanced Photon Source, Argonne National Laboratory. The cryostream temperature was set to 289 K , and data were collected using a Rayonix MX340-HS detector with undulators U23 at 10.74 mm and U27 at 15.85 mm (wavelength range of 1.02 – 1.16 \AA). The beam size was approximately $90 \mu\text{m}$ (h) \times $60 \mu\text{m}$ (v) and slit settings were $200 \mu\text{m}$ (h) \times $70 \mu\text{m}$ (v). Data collection proceeded in four 180° passes with 4° , 4° , 2° and 1° steps, respectively, and with matching offsets to maximize coverage of reciprocal space (Extended Data Fig. 3 and Supplementary Table 2). Laue data were processed by using Precognition and EpiNorm software, with concurrent processing of OFF and ON frames. The data were integrated to 1.8 \AA (Supplementary Table 3) and merged in space group $P1$ using the $C2$ unit cell dimensions. The orientation of the imposed electric field relative to the crystal lattice was established directly from indexed diffraction patterns.

Data for the high-resolution room-temperature (277 K) structure (Fig. 4g and Extended Data Table 3) were collected at the Stanford Synchrotron Radiation Lightsource (SSRL, 11-1) using the PILATUS 6M PAD detector from a single crystal and indexed, integrated, scaled and merged in HKL2000 (ref. 41) (HKL Research). The data showed little radiation damage (HKL2000 radiation-damage coefficients of 0.01 – 0.03 ; values >0.1 – 0.15 indicate significant damage⁴²) or non-isomorphism (coefficient 0.001).

Refinement (C2 OFF models). We refined the structure of LNX2^{PDZ2} in the absence of electric field (OFF) first using the high-resolution (1.1 \AA) data set collected at SSRL at 277 K , with initial phases obtained by molecular replacement using a cryo structure of LNX2^{PDZ2} (model PDB accession 2VWR). After early simulated annealing, a model was refined by alternating rounds of automated refinement in PHENIX⁴³ and manual adjustments in Coot⁴⁴. Alternate conformations were placed where supported by averaged kick⁴⁵ and $F_o - F_c$ maps. The final model had no Ramachandran outliers. Further refinement yielded a model without alternate conformations, also without Ramachandran outliers (Extended Data Table 3). Initial phases for the 289 K OFF data set collected at BioCARS were determined by direct placement of the high-resolution single-conformer model of LNX2^{PDZ}, with small differences in unit cell dimensions refined by rigid-body refinement in PHENIX. Solvent molecules and alternate conformations were modelled in Coot, with real-space refinement to relieve backbone strain, and limited additional refinement in PHENIX (Extended Data Table 2). Anisotropic displacement parameters were refined only for residues with substantial difference density at atomic positions. Note that for calculation of internal difference maps, it is essential that the model used for phasing be refined in the space group of the OFF crystal lattice to guarantee exact position of symmetry elements. We subsequently expanded the refined model to the asymmetric unit of the reduced-symmetry space group using PDBSET (CCP4 6.4.0)⁴⁶.

Internal difference maps. Difference map Fourier coefficients were calculated directly from merged structure factors using custom MATLAB (Mathworks Inc.) scripts performing the following operations: (1) match structure factors F_{hkl} and $F_{\bar{h}\bar{k}\bar{l}}$ and calculate differences $\Delta F_{hkl} = F_{hkl} - \gamma_{\bar{h}\bar{k}\bar{l}} F_{\bar{h}\bar{k}\bar{l}}$, where $\gamma_{\bar{h}\bar{k}\bar{l}}$ are the correction coefficients for absorption anisotropy derived from OFF data (below); (2) obtain phases of the corresponding structure factors from the $C2$ OFF model expanded into $C1$ using PDBSET (CCP4 6.4.0); (3) calculate weights according to

$$w_{hkl} = \left[1 + \frac{\sigma^2(\Delta F)}{\langle \sigma^2(\Delta F) \rangle} + 0.05 \frac{|\Delta F|^2}{\langle |\Delta F|^2 \rangle} \right]^{-1}$$

as previously described^{29,47} and modified⁴⁸ to include a term reducing the contribution of any single structure factor difference. Following Schmidt *et al.*^{29,31}, the difference density maps are improved if structure factors corresponding to large lattice spacings are rejected (here $d_{hkl} > 4 \text{ \AA}$), since EF-X typically produces small-scale electron density differences. For anisotropic absorption correction, we compute $\gamma_{\bar{h}\bar{k}\bar{l}} = \tilde{F}_{\bar{h}\bar{k}\bar{l}}^{\text{OFF}} / F_{\bar{h}\bar{k}\bar{l}}^{\text{OFF}}$, where the tilde indicates local scaling⁴⁹ as implemented in SOLVE⁵⁰. Map coefficients were calculated in PHENIX (FFT) with a grid spacing of 0.3 \AA . Absolute difference density was integrated in UCSF Chimera²⁷, with calculations based on the $C2$ OFF model, expanded to $C1$.

Refinement of excited states. Since the EF breaks $C2$ symmetry, refinement of the up and down models was carried out in the $P1$ space group, with the $C2$ OFF model as a starting point. A $P1$ unit cell was chosen containing one up and one down chain, requiring a rotation around the c^* axis by $\arctan(b/a)$ (here: 31.1°). To do

this, the C2 OFF model was 'expanded' in PDBSET (CCP4 4.6.0) using symmetry operations (1) X, Y, Z , and (2) $(1 - X), Y, (1 - Z)$ and the resulting model was rotated in PDBSET by the specified angle.

Extrapolated structure factors were calculated as $F^{\text{ESF}} = N(F_0^{\text{ON}} - F_0^{\text{OFF}}) + F_0^{\text{OFF}}$, where $N = 1/(1 - f)$ is the extrapolation factor³⁰. For traditional pump-probe experiments, f is interpreted as the fraction of molecules excited by an optical pulse³⁰; here, it increases the effective population of excited states, facilitating structure refinement. N was chosen as a trade-off between two criteria: map quality, which deteriorates with increasing N , and the appearance of difference electron density peaks consistent with internal difference maps, which initially increases with increasing N (systematic optimization of N in a site-specific manner will be explored in future work). Refinement was performed mostly manually in Coot with determination of R factors in PHENIX, combined with bulk solvent scaling and occupancy refinement every 5–10 modifications (Extended Data Fig. 5a). Near completion, a few rounds of overall coordinate refinement (PHENIX, 10–15 microcycles, small geometric weights) were included. Electron density maps and composite omit maps were calculated in PHENIX with 0.3 Å grid spacing and default settings. Reflections in the R_{free} test set were included in final map calculations. The refinement statistics are given in Extended Data Table 2.

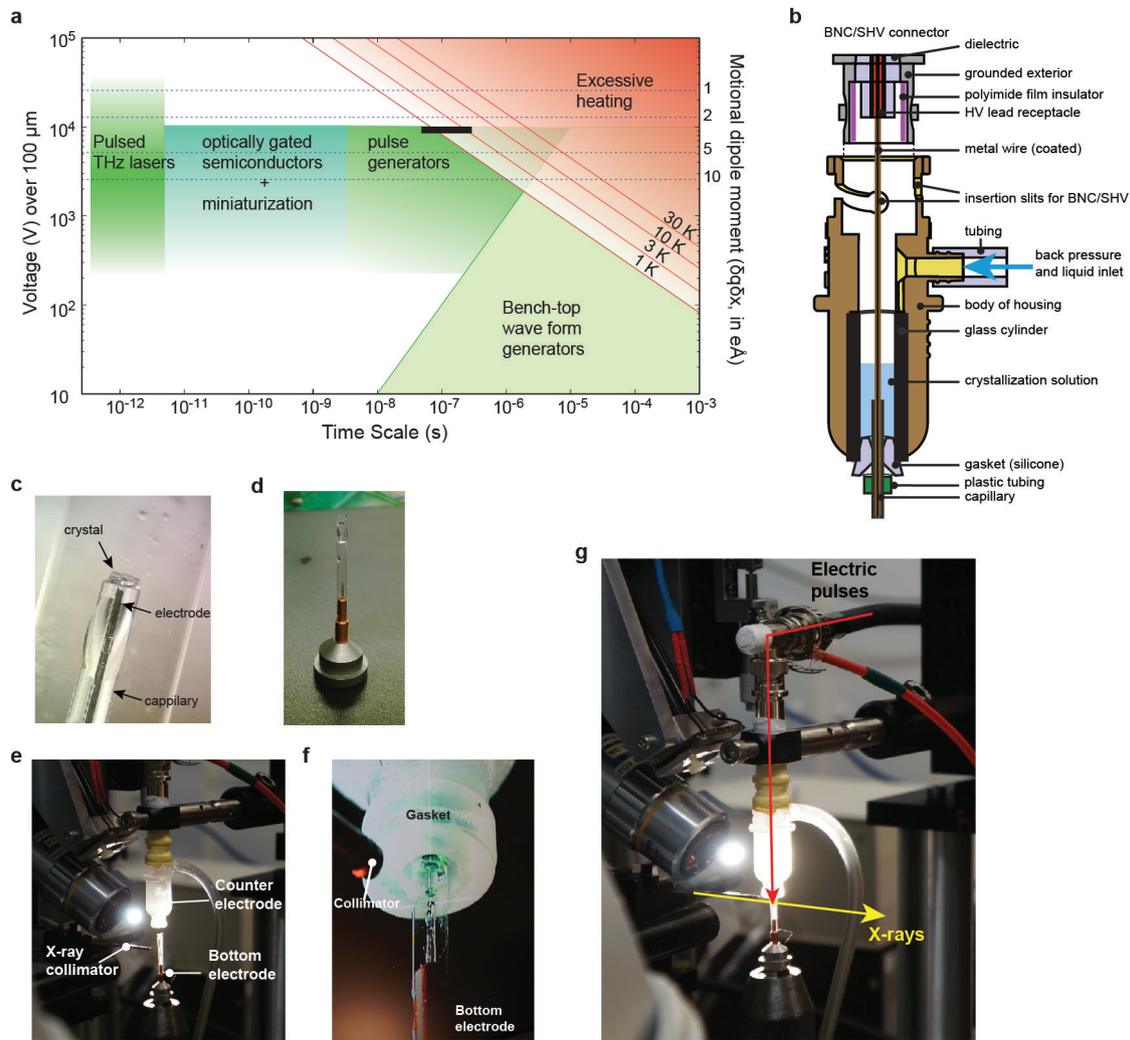
Comparison to homologous PDZ domains. Eleven pairs of high-resolution (≤ 2 Å) X-ray structures of PDZ domains with and without ligand were selected: NHERF-1^{PDZ1}: PDB accessions 1G9O, 1GQ4; PALS-1^{PDZ}: 4UU6, 4UU5; Tiam-1^{PDZ}: 3KZD, 4GVC; ZO-1^{PDZ1}: 4OEO, 4OEP; Erbin^{PDZ}: 2H3L, 1MFG; Dishevelled^{PDZ}: 2F0A, 1L6O; PDZK-1^{PDZ3}: 3R68, 3R69; Shank^{PDZ}: 1Q3O, 1Q3P; GRIP-1^{PDZ6}: 1N7E, 1N7F; PTP-1E^{PDZ2}: 3LNX, 3LNY; PSD-95^{PDZ3} (R.R. *et al.*, unpublished observations). Structures were aligned in PyMOL, using 'super' for backbone atoms, first to the down state of LNX2^{PDZ2}, and then within each pair (Extended Data Fig. 5d legend). For backbone atoms with matching positions in LNX2^{PDZ2}, displacements (Δr) from unbound (apo) to bound (liganded) were then calculated. Atoms with $|\Delta r| < 0.1$ Å were excluded from analysis. Average displacements displayed in Fig. 5c represent the median magnitude and average direction of apo to liganded displacement over homologues.

Statistics. To assess statistical significance of correlations between various experimental measures, the observed quantities were transformed to stabilize variance, reduce kurtosis and approximate a normal distribution. IADDDAT values (Fig. 3) were square-root transformed, and B -factors (Fig. 3) and displacements (Fig. 5) were log-transformed. To assess the statistical significance of correlations, sample correlation coefficients were then Fisher Z -transformed, and tested for deviation

from a standard normal distribution. For the statistical comparison of the data in Fig. 5b and e, individual residues can be considered independent, yielding $P < 0.001$. More conservatively, one can also take the shorter of the correlation length scales of B -factors and observed displacements (~ 2 residues) as a measure of internal data dependence. This yields a reduced number of independent samples and $P < 0.01$. Thus, the association of Fig. 5b and e is robust to local internal correlations in the data. No statistical methods were used to predetermine sample size.

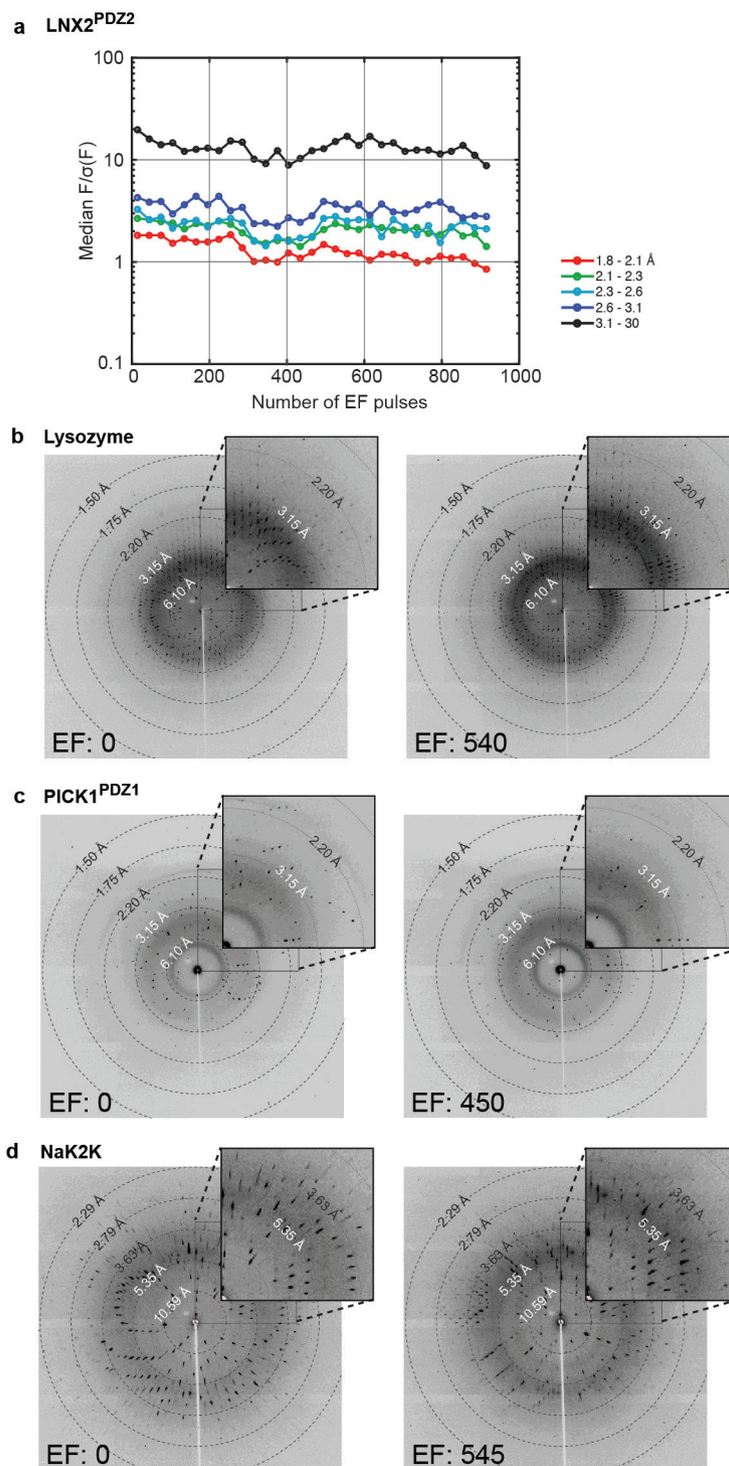
Data availability. Structure factors and refined models have been deposited in the PDB under accessions 5E11, 5E1Y, 5E21 and 5E22.

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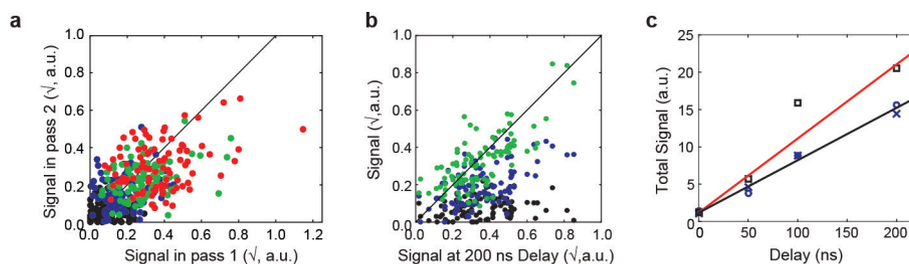
Extended Data Figure 1 | The experimental setup of EF-X. **a**, A plot relating the applied voltage across a 100- μm -thick crystal (left axis) and the size of transition dipole moments of conformational changes that can be excited by $k_B T$ (right axis) to the duration of the applied electric field. Feasible methods of generating strong electric field pulses are indicated as green and cyan shaded areas. Waveform and pulse generators can provide pulses down to the nanosecond timescale. Faster pulses can be generated using terahertz pulsed lasers with strong electric field components⁵¹ or by optical gating of semiconductors⁵²; such systems are already present at third-generation synchrotron and X-ray free-electron laser facilities. The black bar indicates the approximate range covered by the current experiments. The calculation of temperature jumps caused by the electric field is described in Supplementary Information IA. **b**, Schematic

cross-section of the counter electrode. The blue arrow indicates the path by which backpressure is applied to drive flow through the capillary (see Methods). **c**, Crystals are mounted on top of capillaries containing a metal electrode and soaked in crystallization solution. **d**, The capillary with crystal is mounted in a reusable goniometer base and protected from humidity fluctuations with a polyester sleeve (MiTeGen) containing 50% (v/v) crystallization solution. This assembly forms the bottom electrode. **e**, The counter and bottom electrodes are assembled at the beam line to allow rotation around the capillary axis. **f**, Once the sleeve is trimmed to just above the level of the crystal, the counter electrode is brought in using a translation stage (camera view of the approach) (Supplementary Video 2). **g**, Overview of the final set up with the direction of the X-ray and electric field pulses, reproduced from Fig. 1e.



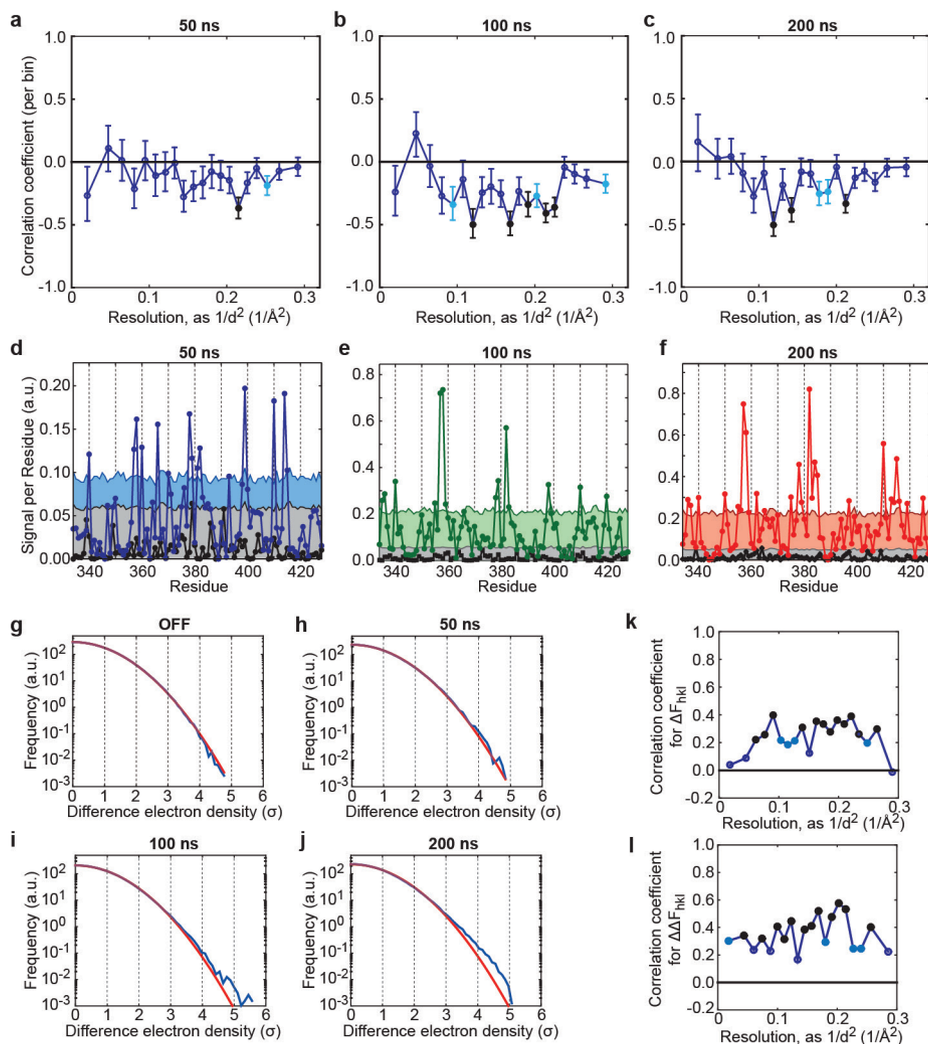
Extended Data Figure 2 | Tolerance of electric field pulses in several protein crystals. **a**, Diffraction quality of a LNX2^{PDZ2} crystal (experiment 3-35, Supplementary Table 2), measured by the ratio of structure factor amplitude to noise ($F/\sigma(F)$) as a function of number of 250 ns, 6 kV electric field pulses and as a function of resolution bin (in colours, see legend). **b-d**, Diffraction images for three other protein crystals before (left) and

after (right) \sim 500 electric field pulses (precise value indicated). Crystal orientations are different between before and after frames. The data correspond to the following experiments in Supplementary Table 2: **b**, lysozyme, experiment 3-08; **c**, PDZ1 of PICK1, experiment 3-17; **d**, NaK2K, experiment 3-80. The data indicate that several protein crystals can tolerate the EF-X experiment.



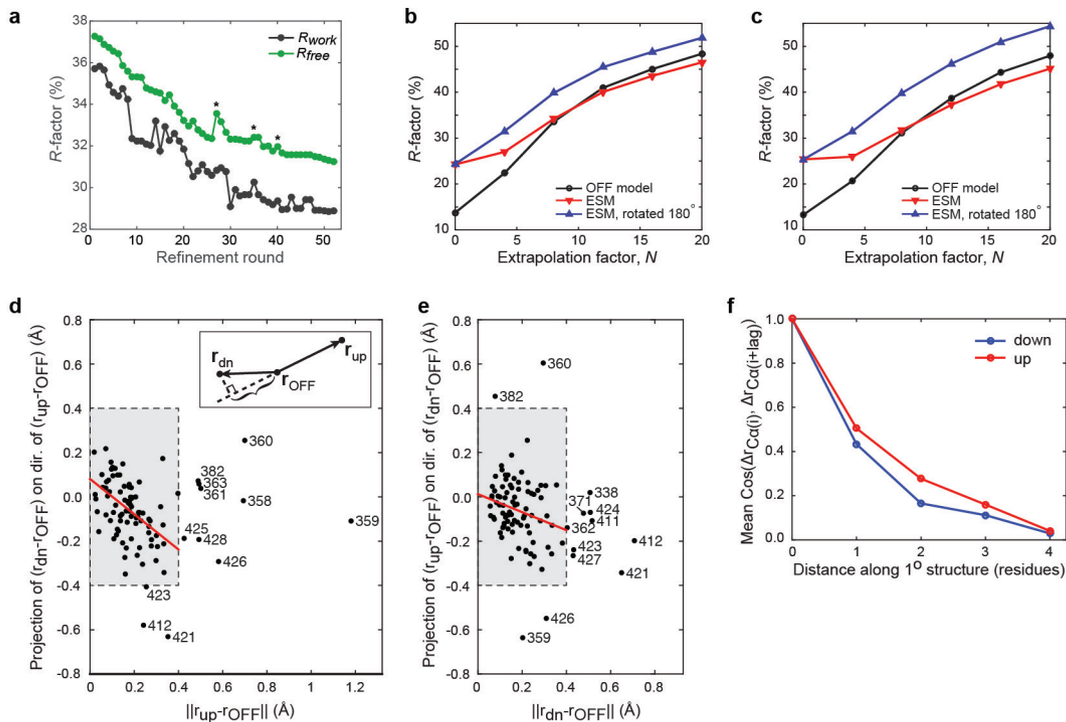
Extended Data Figure 3 | Internal consistency and temporal evolution of internal difference map signal. a–c, Analysis for the data presented in Fig. 3. a, Consistency of estimated signal per residue derived from two data collection passes on the same crystal (black: OFF; blue: 50 ns; green: 100 ns; red: 200 ns). Overall correlation coefficient 0.59. Signal is defined as the integrated absolute difference density above $2.5\sigma_{\text{OFF}}$ within 1.5 \AA of the protein backbone, square-root transformed to stabilize variance. Per-time-point correlation coefficients are: -0.07 (OFF, $P > 0.1$), 0.23 (50 ns; $P = 0.01$); 0.35 (100 ns; $P < 10^{-3}$) and 0.34 (200 ns; $P < 10^{-3}$).

b, Consistency of the obtained signal per residue between time points. Correlation coefficients are: 0.17 (OFF; $P = 0.05$), 0.55 (50 ns; $P = 1 \times 10^{-9}$) and 0.72 (100 ns; $P < 10^{-20}$). The diagonal is shown for reference. Note that slight correlation in the OFF data set may indicate imperfect correction for anisotropic absorption. c, Signal integrated along the entire protein backbone in passes 1 and 2 (blue crosses and circles, respectively) and over the entire data set (squares). The red line indicates a naive expectation of a $\sqrt{2}$ -fold increase in signal-to-noise ratio.



Extended Data Figure 4 | Validation of signal in structure factors and difference maps. **a–c**, A negative correlation between $\Delta F_{hkl} = \Delta F_{hkl}^{\text{ON}} - \Delta F_{hkl}^{\text{OFF}}$ and $\Delta F_{\bar{h}\bar{k}\bar{l}} = \Delta F_{\bar{h}\bar{k}\bar{l}}^{\text{ON}} - \Delta F_{\bar{h}\bar{k}\bar{l}}^{\text{OFF}}$ is consistent with oppositely directed motions in the up and down states. Analysis is performed over 20 resolution bins to allow for statistical testing. Shown are the correlation coefficients per bin between ΔF_{hkl} and $\Delta F_{\bar{h}\bar{k}\bar{l}}$. In a linear response approximation and in the absence of measurement error, we expect $\Delta F_{hkl} = -\Delta F_{\bar{h}\bar{k}\bar{l}}$. Reflections with $|\Delta F_{hkl}| < \sigma(\Delta F_{hkl})$, $|\Delta F_{\bar{h}\bar{k}\bar{l}}| < \sigma(\Delta F_{\bar{h}\bar{k}\bar{l}})$ or $|\Delta F_{hkl}| > 10$ were excluded from analysis, and likewise for $\Delta F_{\bar{h}\bar{k}\bar{l}}$. Results at 50 ns (**a**), 100 ns (**b**) and 200 ns (**c**). To assess significance, each bin was considered statistically homogeneous, with observations considered independent. Bins with significant negative deviation from 0 (after Fisher Z-transform) are indicated as filled circles ($P < 10^{-3}$: black; $P < 10^{-2}$: light blue). Error bars indicate standard errors based on the assumption of a normal distribution after Fisher Z-transform. **d–f**, Statistical significance of Fig. 3g. Comparison of integrated absolute difference density above 2.5σ , within 1.5 \AA of backbone C, N and O atoms ('signal'; see also Fig. 3a). **d**, Comparison of signal in the OFF state and at 50 ns. The grey-shaded area indicates the 0–95th percentile for random sampling from the OFF map at the same probe volume (because the conformation of the backbone changes from residue to residue, the effective probe volume varies along the protein backbone) and threshold. The blue-shaded area indicates the 0–95th percentile for random sampling using the conservative sampling protocol described in test 4 of the statistical validation. **e**, Same analysis at 100 ns. **f**, Same analysis at 200 ns. Note that we were unable to scale all diffraction images at once, and instead scaled the OFF data with each time point separately. We compare each ON data set to the OFF data as scaled with that time point. As a result, there are small differences between the OFF traces in **d–f**.

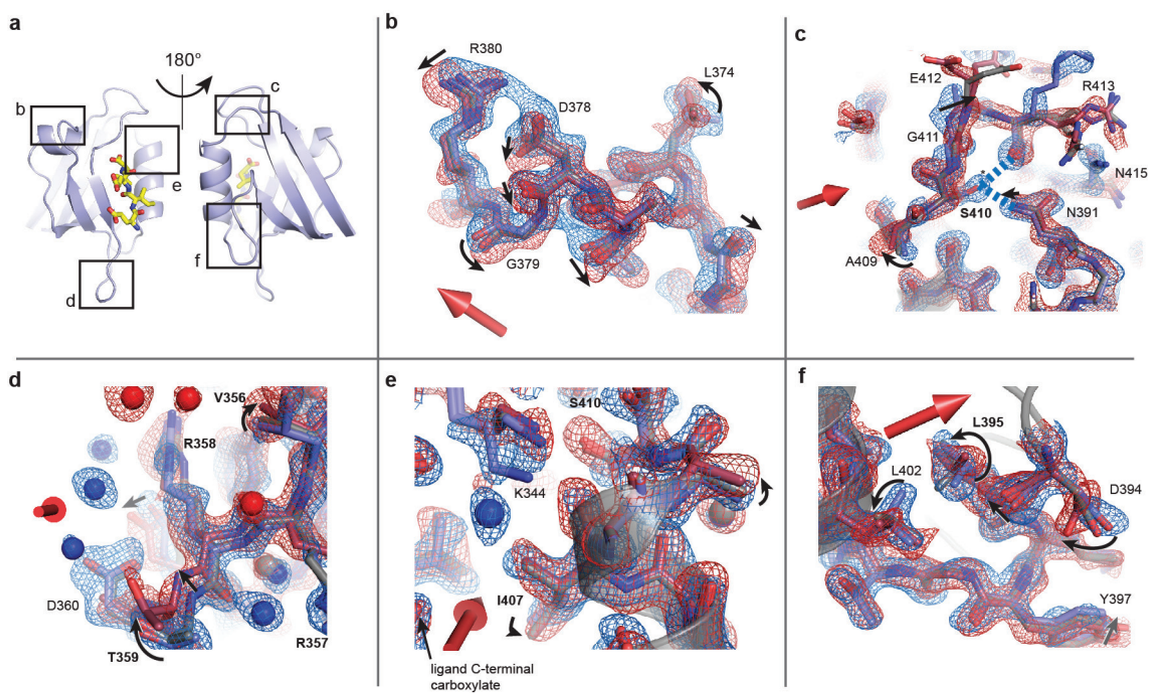
g–j, Deviations from a normal distribution for internal difference maps. Shown are voxel histograms for internal difference electron density (DED) maps without applied field (OFF) (**g**), and at 50 ns (**h**), 100 ns (**i**) and 200 ns (**j**) of applied electric field. Red lines indicate fits to a normal distribution based on calculated variance. Blue lines are histograms of voxel internal DED values (map grids of 0.3 \AA). Note that by construction, for internal DED maps the positive and negative sides of the histogram are the same, apart from discretization effects. To assess statistical significance of deviations from normality, we sampled C2 asymmetric units (ASUs) at the Nyquist sampling frequency (here, 0.9 \AA). For the OFF map, we find no significant deviations from normality ($P > 0.1$ for the Jarque–Bera test, the Anderson–Darling test, and the Lilliefors test; all using default settings in Matlab). At 200 ns, each test rejects a normal distribution with $P < 0.01$. At 50 and 100 ns, the results of statistical testing depend on how the internal DED map is subsampled: for a single C2 ASU, none of the tests rejects the null hypothesis, but when the same number of points is sampled from two neighbouring ASUs, the Jarque–Bera test rejects normality ($P < 0.01$ at 50 and 100 ns), suggesting limited deviation from normality. **k, l**, Reproducibility of a structural response to electric field. Correlation of data set 2 (see Supplementary Table 7) to the data set presented in the text. On the basis of ordinary differences ΔF_{hkl} (**k**) and internal differences $\Delta\Delta F_{hkl} = (F_{hkl}^{\text{ON}} - F_{\bar{h}\bar{k}\bar{l}}^{\text{ON}}) - (F_{hkl}^{\text{OFF}} - F_{\bar{h}\bar{k}\bar{l}}^{\text{OFF}})$ (**l**), reflections with $|\Delta F_{hkl}| < \sigma(\Delta F_{hkl})$, $|\Delta F_{\bar{h}\bar{k}\bar{l}}| < \sigma(\Delta F_{\bar{h}\bar{k}\bar{l}})$, or $|\Delta F_{hkl}| > 10$ were excluded from analysis, and likewise for $\Delta F_{\bar{h}\bar{k}\bar{l}}$. The standard error of correlation coefficient estimates is ~ 0.07 in **k** and ~ 0.10 in **l**. Each bin is statistically homogeneous and observations are considered independent. Resolution bins with significant positive deviation from 0 (after Fisher Z-transform) are indicated as filled circles ($P < 10^{-3}$: black; $P < 10^{-2}$: light blue).



Extended Data Figure 5 | Refinement, voltage-ON model at 200 ns.

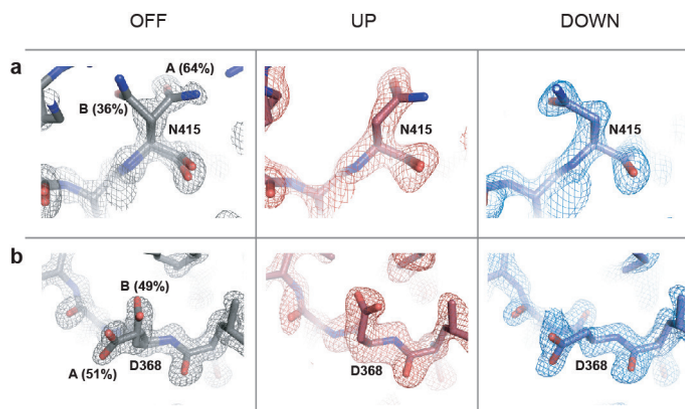
a, Progress of refinement against extrapolated structure factors. Rounds marked by asterisks involved automated refinement with mild stereochemistry constraints to reduce deviations from optimal geometry due to manual refinement in Coot. Fluctuations in R_{work} appear to be mostly due to the PHENIX bulk solvent scaling calculation used in R factor calculation. **b**, R factor for comparison of extrapolated structure factors, as a function of the degree of extrapolation, N , as derived from data set 2 (150 ns; see Supplementary Table 7), against calculated structure factors (F_c) derived from (1) the OFF model (black), (2) the excited state model (ESM) (red), and (3) an 'upside-down' ESM obtained by 180° rotation around the C2 two-fold rotation axis (blue), all derived from data set 1 (Extended Data Table 2). N relates to the fraction f of OFF signal subtracted as $N = 1/(1 - f)$. No refinement against data set 2 was performed except for bulk solvent scaling. No test set was assigned. **c**, For comparison, the same analysis as in **b**, comparing the OFF model and 200 ns ESM model to the 100 ns data (from the same

crystal). **d-f**, Relationship between $C\alpha$ displacements in the up and down conformations at 200 ns. **d**, **e**, Projection of the down displacement on the direction of the up displacement (**d**), and the up displacement on the down displacement direction (all displacements are relative to the OFF model) (**e**); models were superimposed using PyMOL, using C, $C\alpha$ and N atoms of the protein backbone and including only residues 338–356, 362–380, 384–408 and 412–419; this excludes N- and C-terminal regions and mobile parts of the β_2 – β_3 , α_1 – β_4 and α_2 – β_6 loops). Shown are, for example $\Delta r^{\text{down}} \cdot \Delta r^{\text{up}} / |\Delta r^{\text{up}}|$ versus $|\Delta r^{\text{up}}|$, as illustrated in the inset. For small displacements, a simple inverse dependence is expected. This is tested by robust linear regression for (projected) displacements smaller than 0.4 \AA (red line fits to data in grey boxes; using default settings in Matlab). **d**, Slope = -0.80 ± 0.16 , intercept = $0.081 \pm 0.031 \text{ \AA}$; correlation coefficient: -0.44 . **e**, Slope = -0.41 ± 0.17 , intercept = $0.012 \pm 0.033 \text{ \AA}$; correlation coefficient: -0.27 . **f**, Average cosine between displacements of nearby $C\alpha$ atoms as a function of distance along the primary structure.



Extended Data Figure 6 | Additional views of conformational changes due to the electric field. **a**, Reference model indicating regions examined in **b–f**. **b–f**, Maps and models as in Fig. 4, with motions indicated by arrows and residues coupled to ligand binding in PDZ domains shown (as in Supplementary Table 1). **b**, Top view of the α_1 helix, waters omitted and the side chain of Q377 truncated for clarity. **c**, Transverse shift of the α_2 - β_6 loop, and perturbed down state of S410, forming new hydrogen bonds to

R413 and N391 (dashed blue lines). **d**, Upward motion of the β_2 - β_3 loop and change in dynamic disorder of protein and solvent. **e**, Conformational changes at the top of the ligand-binding pocket, with motion of the terminal amine of the K344 towards the ligand carboxylate group in the down state. **f**, Coupled rotameric changes of L402 (α_2 helix), L395 and D394.



Extended Data Figure 7 | Biasing pre-existing conformational heterogeneity in the LNX2^{PDZ2} ground state structure by the external electric field: additional examples. **a, b**, A high-resolution (1.1 Å) room-temperature structure of the voltage-OFF ground state of LNX2^{PDZ2} (Extended Data Table 3), shows partial occupancy of N415 (**a**) and D368 (**b**) in two rotameric states (left). This pre-existing conformational equilibrium is biased in the presence of the electric field (6 kV, 200 ns delay), such that the up and down models each adopt one of the two ground state configurations (middle and right). This supports the result shown in Fig. 4g.

Extended Data Table 1 | Estimates of dipole moments associated with conformational changes

Conformational change	Transition dipole moment (eÅ)	Electric field required for $1 k_B T$ bias (MV/cm)
180° flip of a water molecule ⁴⁶	0.8	3.3
180° flip of a peptide bond ⁴⁶	1.5	1.7
Rotamer flip of a protonated histidine	5.0	0.5
20° turn of a 3-turn α helix dipole ⁴⁶	5.3	0.5
2-ion hop in the KcsA channel ⁴⁹	7.0	0.4
GPCR gating (net, m2R receptor)	~20	~0.13
Gating of a K ⁺ channel ⁵⁰	~100	~0.03

Transition dipole moments were estimated based on the indicated references and for the histidine side chain based on measurements in PyMOL. Shown is the electric field required to bias a conformational equilibrium by $1 k_B T$ with the electric field applied parallel to the transition dipole moment.

Extended Data Table 2 | Data collection and refinement statistics for LNX2^{PDZ2} for EF-X experiment

	LNX2 ^{PDZ2} (OFF)	LNX2 ^{PDZ2} (200 ns)	Extrapolated Differences (8x)
Data collection [‡]			
Space group	C2 [†]	P1	P1
Cell dimensions			
<i>a</i> , <i>b</i> , <i>c</i> (Å)	65.30, 39.45, 39.01	38.15, 38.15, 39.01	38.15, 38.15, 39.01
α , β , γ (°)	90, 117.54, 90	113.31, 113.31, 62.28	113.31, 113.31, 62.28
Resolution (Å)	30.08-1.80 (2.0-1.8)*	30.08-1.80 (2.0-1.8)	30.08-1.80 (1.86-1.80)
<i>R</i> _{sym} or <i>R</i> _{merge}	0.088 (0.051)	0.087 (0.053)	n/a
<i>I</i> / σ <i>I</i>	20.7 (37.1)	20.4 (39.9)	6.98 (0.67)
Completeness (%) [‡]	75.1 (42.5)	72.4 (38.1)	70.2 (17.8)
Redundancy	5.8 (3.9)	5.7 (3.6)	n/a
Refinement			
Resolution (Å)	30-1.8 (1.88-1.8)	30-1.8	30-1.8 (1.88-1.8) [§]
No. reflections [¶]	6,565 (288)	11,568	11,291 (288)
<i>R</i> _{work} / <i>R</i> _{free} (%)	13.2/14.8		28.9/31.3
No. atoms (excl. H)	929		1,883
Protein	829		1,712
Ligand/ion	0		6
Water	94		165
B-factors	21.9		16.7
Protein	19.3		16.0
Ligand/ion	n/a		49.5
Water	35.6		22.5
R.m.s deviations			
Bond lengths (Å)	0.020		0.018
Bond angles (°)	1.63		1.80

All data were collected from a single crystal of LNX2^{PDZ2} using Laue crystallography.

*Highest-resolution shell is shown in parentheses. Data reduction in Precognition (Renz Research) differs from conventional data reduction in that weak spots are discarded a priori, resulting in low apparent completeness and high apparent signal and *R*_{merge}, especially at high resolution. Note also that data statistics are reported after global scaling. Subsequent local scaling slightly affects statistics but this scaling mode does not report full last shell statistics. Extrapolated differences were assessed in Xtrriage (PHENIX).

†For the purpose of refinement of the OFF model, P1 reflections were merged according to C2 symmetry (merging *R* factor for this: 0.077).

‡See Supplementary Table 3 for data collection and reduction statistics as reported traditionally for Laue crystallography, including assessment of completeness in both C2 and P1. Reported data collection statistics refer to P1.

§Data were retained to the resolution of the two 'parent' data sets (OFF and 200 ns); effective resolution based on propagation of errors is 2.3 Å; completeness over 30–2.3 Å is 89.9%.

||Test sets comprised 5% and 10% of reflections for refinement of the OFF model and refinement against extrapolated structure factors, respectively.

¶The matching number of reflections in the high-resolution shell is coincidental.

Extended Data Table 3 | Data collection and refinement statistics for LNX2^{PDZ2} by room-temperature crystallography

LNX2 ^{PDZ2} (high-resolution)		
Data collection		
Space group	C2	
Cell dimensions		
<i>a</i> , <i>b</i> , <i>c</i> (Å)	64.91, 39.29, 38.80	
α , β , γ (°)	90, 117.41, 90	
Resolution (Å)	34.45-1.05 (1.05-1.01)*	
<i>R</i> _{sym} or <i>R</i> _{merge}	0.051 (0.54)	
<i>I</i> / σ <i>I</i>	12.84 (0.45)	
Completeness (%) [†]	77.6 (3.0)	
Redundancy	5.8 (1.2)	
Refinement		
	With alternate conformations	No alternate conformations
Resolution (Å)	34.45-1.05	34.45-1.05
No. reflections	35,251 (137)	35,251 (137)
<i>R</i> _{work} / <i>R</i> _{free}	11.9/13.4	12.6/14.0
No. atoms (non-H)	1,039	824
Protein	929	719
Ligand/ion	0	0
Water	104	99
B-factors	19.2	19.7
Protein	17.1	17.3
Ligand/ion	n/a	n/a
Water	37.1	36.2
R.m.s deviations		
Bond lengths (Å)	0.022	0.023
Bond angles (°)	1.86	1.88

On the basis of data collected from a single crystal.

*Highest-resolution shell is shown in parentheses. Data were retained based on CC1/2 > 50%. *I*/ σ *I* falls below 2 at 1.08 Å.

†Completeness over 50–1.5 Å is 98.2%. Completeness falls below 50% (*I*/ σ *I* = 1) at 1.1 Å.