Elastoresistivity in the incommensurate charge density wave phase of BaNi$_2$(As$_{1-x}$P$_x$)$_2$

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Electronic nematicity, the breaking of the crystal lattice rotational symmetry by the electronic fluid, is a fascinating quantum state of matter. In this work, using electronic transport under strain we investigate the electronic nematicity of BaNi$_2$(As$_{1-x}$P$_x$)$_2$, a candidate system for charge-induced nematicity. We report a large $B_{1g}$ elastoresistance coefficient that is maximized at the tetragonal-to-orthorhombic transition temperature, that slightly precedes the first-order triclinic transition. An hysteretic behavior is observed in the resistance versus strain sweeps and interpreted as the pinning of orthorhombic domains. Remarkably, the elastoresistance only onsets together with a strong enhancement of the incommensurate charge density wave of the material, strongly suggesting that this electronic instability is uniaxial in nature and drive the orthorhombic transition. The absence of sizeable elastoresistance above this electronic phase clearly contrasts dynamic and static electronic nematicity. Finally, the elastoresistance temperature dependence that strongly differs from the Curie-Weiss form of iron-based superconductors reveals major differences for the respective coupling of electronic nematicity to the lattice. Our results uncover an extremely strain-sensitive platform to study electronic anisotropy induced by a charge-density-wave instability.

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INTRODUCTION

With the discovery of iron-based superconductors$^1$, electronic nematicity has emerged as a potential key ingredient for high-temperature superconductivity. Indeed, the observation of strong electronic nematic fluctuations at the optimal conditions for superconductivity suggests that such fluctuations might promote higher $T_c$.$^2$–$^6$. This view is supported by reports for nematicity in other unconventional superconductors as heavy fermions$^7$–$^8$ or cuprates$^9$–$^{13}$. Nonetheless, the best-understood case is by far the one of the iron pnictides, where the electronic nematic fluctuations induce a tetragonal-to-orthorhombic phase transition at $T_c$. The nematic transition, when not coincident with, is closely followed by antiferromagnetic order at $T_N$.$^{14}$–$^{18}$. From this empirical observation and theoretical considerations$^5$ anisotropic magnetic fluctuations are a leading candidate for the mechanism of nematicity in iron-based superconductors. However, beyond the iron pnictides case, and in particular in the absence of long-range magnetic order$^{19}$–$^{24}$, much remains to be understood about the mechanisms of electronic nematicity and its significance for superconductivity.

In this regard, BaNi$_2$As$_2$, which shares the same high temperature tetragonal structure as the intensively studied BaFe$_2$As$_2$, has recently attracted attention. In contrast to its iron-analog, BaNi$_2$As$_2$ is superconducting below $T_c \approx 0.7\, K$ at ambient pressure$^{25}$ and hosts two types of charge density waves (CDWs). Upon cooling, first appears an incommensurate charge density wave (ICDW) that develops strongly at $T \approx 155\, K$. A weaker diffuse signal can be tracked all the way up to room temperature$^{26}$–$^{28}$. The exact nature of this I-CDW is currently under intense investigation. The reciprocal space pattern of the I-CDW superlattice peaks reported by $X$-ray diffraction experiments indicated that the charge modulation is unidirectional$^{29}$–$^{32}$. Thus, it has been initially linked to a breaking of the crystal lattice rotational symmetry$^{33}$–$^{35}$. Although, from a wide reciprocal lattice mapping it has been later suggested that, overall, the I-CDW is a symmetric, biaxial, rotationally invariant state$^{36}$, a clear rotational symmetry breaking has been recently observed within the I-CDW phase by high-resolution thermal expansion measurements$^{26,32}$. At a slightly lower temperature, a commensurate uniaxial charge density wave (C-CDW) develops at the expense of the former$^{26}$–$^{30}$ and is associated with a first-order triclinic structural transition at $T_{\text{tric}} \approx 137\, K$. This latter coincident transition is suppressed through numerous chemical substitutions at a critical value, $x_{\text{cr}}$.$^{25}$–$^{27,29,30,33,34}$

Despite the absence of static magnetism down to the lowest investigated temperature$^{35}$ possible indications for electronic nematicity have been reported. First, in Ba$_{1-x}$Sr$_x$Ni$_2$As$_2$ strain-dependent electrical transport measurements, namely elastoresistance, have been interpreted as a signature of a large $B_{1g}$ electronic nematic susceptibility upon approaching the triclinic phase transition in substituted samples$^{31}$. Second, a continuous orthorhombic transition was recently found to precede the triclinic one in Ba(Ni$_{1-x}$Co$_x$)$_2$As$_2$ and BaNi$_2$S$_2$.$^{32}$. Unlike in BaFe$_2$As$_2$ the in-plane orthorhombic axes are aligned with the tetragonal ones and the associated lattice distortion is much smaller. By analogy with the iron pnictides, it was proposed that this intermediate phase with broken rotational symmetry is a possible manifestation of charge-induced nematicity$^{26}$.

In addition, the superconducting $T_c$ has been found to sharply increase to $\approx 3\, K$ for substitution levels just above the suppression of the triclinic and C-CDW transition, but the origin of this enhancement is still debated. In Ba$_{1-x}$Sr$_x$Ni$_2$As$_2$ it has been associated with electronic nematic fluctuations$^{31}$, while in BaNi$_2$S$_2$ it has been attributed to an enhanced electron-phonon coupling through a lattice softening$^{34}$. Thus, it is critically needed to assess the possible advent of electronic nematicity in these materials and establish its interplay...
with the aforementioned lattice and electronic instabilities. In this work, we investigate the electronic nematicity of BaNi$_2$(As$_{1-x}$P$_x$)$_2$ with $0 \leq x \leq 0.10$, a system for which the tetragonal-to-orthorhombic transition is established with using elastoresistance measurements. We report a large maximum of the $B_{1g}$ symmetric elastoresistance coefficient, $m_{12}$, that occurs, up to $x = 0.075$, at the temperature of the orthorhombic transition, where the rotational symmetry is broken. Importantly, the elastoresistance onset corresponds to a strong increase in the $I$-CDW superlattice peak intensity that cannot be described by a Curie-Weiss-like temperature dependence and points to a minor coupling of electronic nematicity to the lattice. Thus, the anisotropic strain-dependent electrical transport is a property of the $I$-CDW phase as done in the following, the in-plane resistance anisotropy that strongly suggests the latter as being uniaxial in nature. Importantly, dynamic nematic effects recently reported by Raman scattering significantly precede the unveiled static nematic order parameter in the associated electronic nematic susceptibility, consistent with a broad temperature range of dynamic anisotropic charge/orbital correlation prior to the static $I$-CDW. Finally, a careful investigation of the hysteretic behavior of the resistance versus strain sweeps strongly suggests that the hysteresis originates from the pinning of orthorhombic domains.

RESULTS

Experimental details

We start by investigating the freestanding resistance of BaNi$_2$(As$_{1-x}$P$_x$)$_2$ single crystals in Fig. 1a. A metallic behavior with residual-resistivity ratio (RRR) values in line with the literature is observed. For BaNi$_2$As$_2$, $\text{RRR} \approx 12$ i.e. approximately the value found in BaFe$_2$As$_2$. In agreement with previous reports, a sharp increase of electrical resistance occurs at $T_n \approx 137 \text{ K}$ upon cooling in BaNi$_2$As$_2$, signaling the triclinic structural transition. The elastoresistance indicates the first-order nature of the transition. Upon increasing P-substitution this transition is shifted towards lower temperature, the resistance upturn becomes a downturn, and the width of the thermal hysteresis increases. No such transition is observed for $x = 0.10$, where cooling and warming measurements overlap, indicating a critical doping for the triclinic phase $x_c \approx 0.08$, in agreement with the literature.

At low temperature, and except at the highest ($x = 0.10 > x_c$) P-content, the electrical resistance does not follow a $T^2$ temperature dependence (see Supplementary Note 6 for more details). At high enough temperatures the resistance is well described as $R = R_0 + A T^n$ (see dashed black line in Fig. 1a for $x = 0.070$), with $A$ and $n$ being x-dependent. As seen in the inset of Fig. 1a the exponent $n$ shows a significant increase across the triclinic critical point, and in particular $n = 1$, i.e., a linear-in-temperature resistance is observed within a narrow substitution range around $x_c$, whose origin is still unknown (more details on the fitting are given in Supplementary Note 1).

In order to study the elastoresistance of BaNi$_2$(As$_{1-x}$P$_x$)$_2$, we induce an antisymmetric biaxial strain to our single crystals by gluing them on top of a piezo stack as visualized in Fig. 1b, a technique initially used in strongly correlated systems in ref. This method allows to extract the elastoresistance coefficients defined by $m_{ij} = 1/R_i (\partial R_i/\partial \epsilon_{jj})$, where $j$ denotes the direction of the strain and $i = x$ (resp. $i = y$) corresponds to longitudinal (resp. transverse) measurements with respect to the piezo poling axis, $x$. In the following, we use the Voigt notation for the elastoresistance coefficients, in particular $xx = 1$ and $yy = 2$, and formulate in terms of the irreducible representations of the high-temperature $D_{4h}$ tetragonal point group.

With the [100]$_{\text{tet}}$ axis aligned to the piezo stack poling direction, as done in the following, the in-plane resistance anisotropy that develops under strain is proportional to the symmetry-resolved $B_{1g}$ elastoresistance coefficient, $m_{12} - m_{11}$,

$$\frac{\Delta R}{R_{xx}} - \frac{\Delta R}{R_{yy}} = (m_{11} - m_{12}) (\epsilon_{xx} - \epsilon_{yy})$$  \hspace{1cm} (1)

Since for sufficiently small anisotropy any potential electronic nematic order parameter is proportional to the resistance anisotropy, the associated electronic nematic susceptibility in the $B_{1g}$ channel is probed by the $m_{12} - m_{11}$ elastoresistance coefficient.

Elastoresistance of BaNi$_2$As$_2$

A typical example of raw data in BaNi$_2$As$_2$ in the transverse geometry is shown in Fig. 1c. The corresponding detailed temperature dependence of the elastoresistance measurement is reported in Fig. 2.

First, we show the normalized resistances as a function of temperature with the sample being glued to the piezo, see Fig. 2a. While both longitudinal and transverse directions (squares and circles, respectively) overlap at high enough temperature, a clear

![Elastoresistance of BaNi$_2$As$_2$](image)
discrepancy appears below $T \approx 145$ K, that corresponds to the second-order orthorhombic transition temperature, $T_{\text{orth}}$, as determined by thermal expansion measurements\textsuperscript{26,32}. This observation evidences that, within the orthorhombic state, the sample size is at least partially, detwinned through the anisotropic thermal expansion of the piezo (see Supplementary Note 2) and the resulting strain.

The normalized freestanding resistance (dark and light red lines for the warming and cooling measurement, respectively), reproduced from Fig. 1a, also overlaps with that of the sample glued to the piezo down to $\approx 160$ K. However, at lower temperatures, a difference appears which can be attributed to the differential thermal expansion of the piezo and BaNi$_2$As$_2$ together with the emergence of a finite elastoresistance. The similar resistivity values observed in the longitudinal and transverse channels down to $T_{\text{orth}}$ are also in line with the mainly in-plane isotropic strain arising from the thermal expansion mismatch (see details in Supplementary Note 2) and points to a finite in-plane symmetric elastoresistance.

While the gluing induces a broadening of the triclinic transition as seen by electrical transport, a thermal hysteresis is still clearly observed in these strained conditions and the transition temperature is not substantially shifted. Finally, note that the sign of the resistivity anisotropy in the orthorhombic state is similar to the one in BaFe$_2$As$_2$, i.e., the smaller in-plane orthorhombic axis (aligned with the $y$ piezo axis) is the one with the higher resistivity\textsuperscript{26,37}.

The linear slopes of the resistance versus strain sweeps, $1/ R_0(dR_0/dx_0)$, are reported in Fig. 2b. Note that they are extracted during the same temperature cycle as the corresponding resistances of Fig. 2a, upon applying a voltage to the piezo stack at fixed temperatures. At high temperature, no response to strain is seen in any channel, in agreement with the overlap of the respective electrical resistances. However, below $T_{\text{onset}} \approx 160$ K an elastoresistance signal develops sharply in both directions and peaks at $T^* \approx 145 \pm 2$ K $= T_{\text{orth}}$, with an opposite sign along the two directions. Thus, the $B_{1g}$ symmetric $m_{12} - m_{11}$ elastoresistance coefficient extracted from the difference of the longitudinal and transverse measurements is maximum at $T^* = T_{\text{orth}}$ (see Eq. (1) and Fig. 3a; in the following, $T^*$ is formally defined as the temperature of the $m_{12} - m_{11}$ maximum). Notably, no strong feature appears at the triclinic transition temperature. This brings us to our first important result: the $B_{1g} \cdot$ symmetric $m_{12} - m_{11}$ maximum occurs at the orthorhombic transition rather than at the triclinic one, in contrast to what has been previously reported for Sr-substituted samples\textsuperscript{31}. This is fully consistent with the 4-fold symmetry breaking that occurs at the orthorhombic transition\textsuperscript{26,32} and the absence of thermal hysteresis in the elastoresistance response, in particular around $T^*$ (see also additional measurements in Supplementary Fig. 4).

**Evolution with P-concentration**

The $m_{12} - m_{11}$ elastoresistance coefficient at the different P-contents investigated is shown in Fig. 3b–d, as a function of $T^* = T_{\text{orth}}$ where the triclinic transition temperature and the elastoresistance coefficient are extracted upon cooling. A maximum of the elastoresistance is found up to the highest concentration investigated at temperature exceeding the triclinic transition. Rather, up to $x = 0.075$, $T^*$ is in good agreement with $T_{\text{orth}}$ as determined by thermal expansion. With increasing substitution level, $T^*$ decreases smoothly towards $T^* = 50$ K for $x = 0.10$ where the associated elastoresistance maximum becomes weaker and significantly broader. In particular, there is no enhancement of $m_{12} - m_{11}$ associated with the enhanced superconducting $T_c$ in the absence of the triclinic structure for $x = 0.10$. Note that for $x = 0.10$ no orthorhombic distortion is observed by high-resolution thermal expansion, which shows however a clear signature of a different first-order transition\textsuperscript{32}. This is most likely not directly related to the elastoresistance maximum since, for lower P-contents, no evidence for a thermal hysteresis is observed. We discuss this particular case later.

A fundamental aspect is the comparison between the temperature dependences of the $m_{12} - m_{11}$ elastoresistance coefficient and of the I-CDW superlattice peak intensity recorded at $Q_{\text{CDW}} = (4 \pm 0.72)$ $\text{Å}$. Using samples from the same batches across the entire substitution range investigated, we find that both quantities onset at very similar temperatures. The elastoresistance coefficient is negligible above $T_{\text{onset}}$, that corresponds to a strong increase in the I-CDW superlattice peak intensity, but significantly precedes the rotational symmetry breaking at $T_{\text{orth}} < T_{\text{onset}}$\textsuperscript{32}. Thus, a large
This scenario would require measurements under larger strain and is reminiscent of Ba$_1$P$_{11}$Co$_{33}$coef.

No hysteresis is seen, i.e., the response of electrical resistance to P-substitution and temperature. In all samples at high temperature maximum elastoresistance (see arrows). While for BaNi$_2$As$_2$ a peculiar feature of the reported elastoresistance is the presence of reproducible hysteresis in the resistance versus strain sweeps (see Fig. 1c). This was previously reported in Ba$_{1-x}$Sr$_x$Ni$_2$As$_2$ and attributed to the pinning of static nematic domains by the P-concentration and is highest at $x = 0.10$. While this might be an artifact of strain transmission, one possible alternative scenario, that remains to be investigated, is a stronger orthorhombic distortion at this particular substitution content.

**Strain hysteresis**

A peculiar feature of the reported elastoresistance is the presence of reproducible hysteresis in the resistance versus strain sweeps (see Fig. 1c). This was previously reported in Ba$_{1-x}$Sr$_x$Ni$_2$As$_2$ and attributed to the pinning of static nematic domains by the I-CDW ($T_{m1}$, $T_{tri}$) and ascribed to a first-order reorientation of the CDW wavevector with uniaxial stress. In Fig. 4 we report the evolution of the hysteresis width, i.e., the maximum difference between the up and down strain sweeps, shown here for $x = 0.075$ at $T = 70 \, K$ in the transverse geometry. For a given $x$ value, the hysteresis width shown corresponds to the (transverse or longitudinal) measurement with the best signal-to-noise ratio. At any given temperature we report the value obtained from an average of several strain sweeps, the positive (resp. negative) error bars are extracted from the maximal (resp. minimal) value of individual strain sweeps. At temperatures where no hysteresis is resolved, the hysteresis width has to be smaller than the experimental noise and this is shown as a positive error bar extending down to zero. The arrows indicate the corresponding $T$ temperatures of maximum $m_{12} - m_{11}$ (see Fig. 3). For $x = 0.10$ no finite hysteresis is resolved down to the lowest temperature.

For all other substitution levels, we observe a rather sharp increase of the hysteresis width across the $T_{tri}$ temperature of maximum elastoresistance (see arrows). While for BaNi$_2$As$_2$ a finite hysteresis is found to emerge at $T = T_{tri}$, in agreement with ref. 31, this is not the case in P-substituted samples. Since the hysteresis width significantly increases across $T_{tri}$ for $0 \leq x \leq 0.075$ one likely scenario is that the hysteresis is associated to the pinning of orthorhombic domains. This is consistent with the orthorhombic domains orientation. Testing this scenario would require measurements under larger strain and a precise knowledge of the temperature and substitution dependencies of the spontaneous orthorhombic distortion. The remarkable absence of a finite hysteresis at $x = 0.10$ might be a signature of the absence of the orthorhombic distortion, of which the absence of detwinning effect seen in thermal expansion measurements is a more direct evidence. Finally, the hysteretic

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*Fig. 3* Evolution of $m_{12} - m_{11}$ with P-concentration in BaNi$_2$(As$_{1-x}$P$_x$)$_2$. a-d $B_{1g}$ symmetry-resolved $m_{12} - m_{11}$ elastoresistance coefficient (filled circles, left axis) together with the integrated intensity of the I-CDW satellite at $Q_{CDW} = (4.072 \, 1, 0, 0)_{ntr}$ (empty squares, right axis, reproduced from ref. 27), for different $x$ values as indicated. Both quantities are shown upon cooling, except for the $x = 0.10$ X-ray diffraction (XRD) data, and plotted as a function of $T - T_{tri}$ where the triclinic transition temperature is measured upon cooling. No triclinic phase is observed at $x = 0.10$ and thus the data are as a function of $T$ in d. The gray area represents the width of the triclinic transition as determined from the elastoresistance samples (see Supplementary Note 5). Note that elastoresistance and XRD measurements are performed on different samples that share similar triclinic transition temperatures, except for $x = 0.075$ where the XRD sample has a slightly lower P-content and accordingly higher $T_{tri} \approx 55K$. The arrows indicate the onset temperature of $m_{12} - m_{11}$, $T_{onset}$ and the temperature of the elastoresistance maximum, $T_{tri}$. In the inset of a the $B_{2g}$-symmetric $m_{66}$ elastoresistance coefficient of BaFe$_2$As$_2$ from ref. 42 (black diamonds, vertically scaled) is compared to the $m_{12} - m_{11}$ coefficient of BaNi$_2$As$_2$. Lines are guides to the eye.

*Fig. 4* Elastoresistance hysteresis as a function of P-substitution. Temperature dependence of the hysteresis width for the indicated compositions. The inset shows the definition of the hysteresis width, i.e., the maximum difference between the up and down strain sweeps, shown here for $x = 0.075$ at $T = 70 \, K$ in the transverse geometry. For a given $x$ value, the hysteresis width shown corresponds to the (transverse or longitudinal) measurement with the best signal-to-noise ratio. At any given temperature we report the value obtained from an average of several strain sweeps, the positive (resp. negative) error bars are extracted from the maximal (resp. minimal) value of individual strain sweeps. At temperatures where no hysteresis is resolved, the hysteresis width has to be smaller than the experimental noise and this is shown as a positive error bar extending down to zero. The arrows indicate the corresponding $T$ temperatures of maximum $m_{12} - m_{11}$ (see Fig. 3). For $x = 0.10$ no finite hysteresis is resolved down to the lowest temperature.
strain behavior persists into the triclinic phase where the associated structural and/or C-CDW domains probably play a major role.

**DISCUSSION**

We summarize our results in a phase diagram, see Fig. 5. A large $m_{12} − m_{11}$ elastoresistance coefficient onsets together with a strong enhancement of the I-CDW satellite intensity and has a maximum at the orthorhombic transition up to $x = 0.075$. Remarkably, this elastoresistance onset also corresponds to a maximum in the crystallographic $c/a$ ratio which occurs at the $T_{\text{cross}}$ temperature reproduced from ref. 32. At $x = 0.10$, even though no orthorhombic transition has been reported by high-resolution thermal expansion, a weaker and broader $m_{12} − m_{11}$ maximum is found. In parallel, the hysteretic behavior of the elastoresistance reported at lower substitution levels is not resolved anymore. Notably, upon increasing substitution level, the enhanced superconducting $T_c$ above the triclinic critical point coincides with a reduction of $m_{12} − m_{11}$, and not an enhancement. Together with the already $T_c = 3 K$ superconductivity of fully substituted BaNi$_2$P$_2$41, this observation strongly suggests that the superconductivity of BaNi$_2$(As$_{1-x}$P$_x$)$_2$ with $x > x_c$ is not significantly boosted by electronic nematicity.

We now consider in more detail the elastoresistance signal itself. The advent of a large $B_{1g}$ elastoresistance response maximized across the tetragonal-to-orthorhombic transition of BaNi$_2$(As$_{1-x}$P$_x$)$_2$ is certainly reminiscent of the electronic nematic transition of its iron-counterpart, BaFe$_2$As$_2$. However, there are fundamental differences between these two cases, which are highlighted in the following.

In the iron pnictides the $m_{10g}$ elastoresistance coefficient, probing nematicity in the $B_{2g}$ channel, follows a typical Curie-Weiss dependence over a wide temperature range of up to 100 K or more above the orthorhombic transition. Such a temperature dependence is expected on theoretical grounds from the coupling of an electronic nematic order parameter to a lattice strain of the same symmetry2,3,6,7,32. This temperature dependence is also observed above the nematic transition of FeSe$_{1−x}$ whose spin and/or orbital origin is still debated. In sharp contrast, in BaNi$_2$As$_2$, $m_{12} − m_{11}$, probing nematicity in the $B_{1g}$ channel, increases only in the close vicinity of the orthorhombic transition and its onset corresponds to a strong increase in the I-CDW superlattice peak intensity seen in x-ray diffraction experiments. The difference is evident in the comparison shown in the inset of Fig. 3a where BaFe$_2$As$_2$ is chosen as it shares a close orthorhombic transition temperature. While for the latter compound the $m_{11g}$ elastoresistance coefficient can be tracked up to 150 K above the nematic transition, for BaNi$_2$As$_2$ $m_{12} − m_{11}$ $< 1$ only 20 K above $T_{\text{orth}}$. Hence, in BaNi$_2$As$_2$, the temperature dependence of the elastoresistance is very different from that of the well-established electronic nematic systems, and in particular points to a minor coupling of electronic nematicity to the lattice. Moreover, while in the iron-based compound only the rotational symmetry is broken at $T_{\text{orth}}$ in the nickel-based compound this may coincide with the I-CDW transversal symmetry breaking. In this latter scenario, there is no static electronic nematic order in BaNi$_2$As$_2$ in a strict sense since rotational and translational symmetries are broken simultaneously.

With increasing substitution level, the $m_{12} − m_{11}$ maximum broadens, which can be ascribed to the broadening of the orthorhombic distortion seen in thermal expansion, the disorder, and finally, the effect of external stress applied across the orthorhombic transition from the thermal expansion mismatch with the piezo. The latter effect is significant when the externally applied strain is of the same order as the spontaneous orthorhombic distortion. We expect this situation to be realized in BaNi$_2$(As$_{1-x}$P$_x$)$_2$ where, within our experimental conditions, both quantities are $\approx 10^{−3}$ to $10^{−4}$ (see thermal expansion measurements in Supplementary Note 2). In addition, as for $x = 0$, the $m_{12} − m_{11}$ elastoresistance coefficient onsets together with a strong increase in the I-CDW satellite intensity and grows upon approaching the rotational symmetry breaking, advocating for a similar origin. Moreover, even in substituted samples, a Curie-Weiss susceptibility cannot fairly describe $m_{12} − m_{11}$ over a significant temperature range (more details in Supplementary Note 7). Thus, in pure and P-substituted samples, the large $m_{12} − m_{11}$ elastoresistance coefficient is not an evidence for critical electronic nematicity, but is a property of the I-CDW phase. We note that the absence of a significant critical electronic nematicity is consistent with Young modulus measurements at $x = 0.10$ and an ARPES study of BaNi$_2$As$_2$ under uniaxial stress. Finally, it is noteworthy to mention that the elastoresistance onsets at temperature significantly lower than the broadening and subsequent splitting of the $E_{2g}$ phonon mode seen by Raman scattering and ascribed to dynamic electronic nematicity. This difference points to a broad temperature range of dynamic anisotropic charge/orbital correlation preceding the static I-CDW order, a conclusion which is supported by recent inelastic x-ray scattering experiments.

Let us now consider in more detail the parallel strong increase of the I-CDW superlattice peak intensity and the onset of the $m_{12} − m_{11}$ elastoresistance coefficient. Assuming that the temperature dependence of the electron-phonon coupling does not play a pivotal role, as suggested by several works, this observation points to a direct connection between the I-CDW and the $B_{1g}$ elastoresistance.

Regardless of the exact mechanism of formation of the I-CDW, perturbation of the charge-ordered state can be achieved when strain of the right symmetry is applied and a corresponding electrical transport signature is expected through, for instance, modification of the Fermi surface. Consequently, our observation of a large $m_{12} − m_{11}$ elastoresistance coefficient only together with a strong enhancement of the I-CDW satellite intensity points towards a coupling between the I-CDW order parameter, the associated fluctuations, and the $c_{xy}−c_{yx}$ strain, that strongly advocates for an uniaxial nature of the I-CDW. In particular, no
significant $B_{1g}$ elastoresistance is expected to result from a symmetric biaxial I-CDW with very similar modulations along and perpendicular to the tetragonal $a$-axis, as $c_{xy}$ and $c_{yx}$ strains should perturb the I-CDW in an identical manner. In turn, this observation strongly suggests an intimate relationship of the I-CDW with the orthorhombic distortion, that manifests itself as a $m_{12} - m_{11}$ maximum, and that we interpret as a signature of the long-range order of the I-CDW. Importantly, this contrasts with Ba$_{1−x}$Sr$_{x}$Ni$_2$As$_2$ where, for $x > 0.5$, a large $m_{12} - m_{11}$ elastoresistance coefficient occurs in the absence of the I-CDW phase and where, additionally, the fate of the orthorhombic phase remains to be investigated.$^{29,31}$ Our results, in particular the significant sensitivity of the I-CDW to uniaxial stress, put strong constraints on the theoretical description of this phase. It should motivate a refinement of its real space structure and spectroscopic studies under uniaxial stress. Another promising avenue is the recently developed in-situ combination of strain-dependent x-ray diffraction and electrical transport measurements.$^{39}$ Further elastoresistance studies in CDW systems, where a Curie-Weiss dependence is generally not observed$^{39,50}$ are also highly desirable to investigate the different mechanisms at play.

Finally, we focus on a particularly intriguing case, BaNi$_2$(As$_{0.9}$P$_{0.1}$)$_2$, located above the triclinic critical point, $x_c$. As for lower substitution levels, the onset of $m_{12} - m_{11}$ does correspond to a strong increase in the I-CDW superlattice peak intensity. Thus, it is very likely that, as for lower $P$-contents, the $B_{1g}$ elastoresistance reflects the growth of the I-CDW. However, as shown in Fig. 4 a strain hysteresis is not resolved at this substitution content anymore. In parallel, no detwinning of the orthorhombic domains is seen in high-resolution thermal expansion measurements, questioning the occurrence of an orthorhombic phase transition for this composition. A first-order transition is observed in thermal expansion within the temperature range of the $m_{12} - m_{11}$ maximum,$^{32}$ but the absence of a thermal hysteresis in the elastoresistance points toward a different origin. Moreover, the resistance measurements do not reveal any evidence of a phase transition in the vicinity of $T \approx 50$ K (see Supplementary Fig. 7). Noteworthy, the onset of the $m_{12} - m_{11}$ elastoresistance coefficient coincides with a lattice softening as seen by Young modulus measurements, which is argued to be incompatible with critical electronic nematicity.$^{7}$ The simultaneous maximum of elastoresistance and saturation of the softening below $\approx 50$ K strongly suggests a close relationship between those two. The $m_{12} - m_{11}$ maximum, though broad, in the (likely) absence of an orthorhombic transition calls for further investigations.

In conclusion, we report that the I-CDW instability of BaNi$_2$(As$_{1−x}$P$_x$)$_2$ is extremely sensitive to antisymmetric biaxial strain. Specifically, we observe a large $B_{1g}$-symmetric elastoresistance that strongly advocates for an I-CDW uniaxial in nature. Although the elastoresistance maximum at the tetragonal-to-orthorhombic transition is certainly reminiscent of the iron pnictides, the temperature dependence of $m_{12} - m_{11}$ is unprecedented and strikingly contrasts with known examples of electronic nematic transitions. In particular, the absence of a Curie-Weiss-like form points to a smaller coupling of electronic nematicity to the lattice as compared to the iron-based superconductors. The absence of significant $B_{1g}$ elastoresistance above $T_{\text{onset}}$, where a pronounced $E_{1g}$ phonon broadening is observed by Raman scattering, clearly contrasts the static and dynamic electronic nematicity. Importantly, the weakening of the $m_{12} - m_{11}$ elastoresistance coefficient observed in parallel to the enhanced superconducting $T_c$ in the absence of the triclinic structure evidence that the strain-sensitive electronic correlations revealed by elastoresistance are not responsible for the stronger superconductivity. Finally, a careful inspection of the hysteretic behavior of the resistance versus strain sweeps points to the pinning of orthorhombic domains as a likely origin.

METHODS

Single crystals growth and chemical analysis

Single crystals of BaNi$_2$(As$_{1−x}$P$_x$)$_2$ (with $x = 0.0, 0.035, 0.07, 0.075, 0.10$) were grown using a self-flux method. NiAs binary was synthesized by mixing the pure elements Ni (powder, Alfa Aesar 99.9999%) and As (lumps, Alfa Aesar 99.9999%) that were ground and sealed in a fused silica tube and annealed for 20 h at 730 °C. All sample preparations were performed in an argon glove box (O$_2$ content <0.5 ppm). For the growth of BaNi$_2$(As$_{1−x}$P$_x$)$_2$, a ratio of BaNiAs:NiP = 1:4(1−x):4x was placed in an alumina tube, which was sealed in an evacuated quartz ampule (i.e., $10^{-7}$ mbar). The mixtures were heated to 500–700 °C for 10 h, following by heating slowly to a temperature of 1100–1180 °C, soaked for 5 h, and subsequently cooled to 1000–900 °C at the rate of 0.5 °C/h to 2 °C/h, depending on the phosphorus content used for the growth. At 1000–900 °C, the furnace was canted to remove the excess flux, followed by furnace cooling. Plate-like single crystals with typical sizes 3 x 2 x 0.5 mm$^3$ were easily removed from the remaining ingot. The crystals were brittle having shiny brass-yellow metallic luster. Electron micro probe analysis of the BaNi$_2$(As$_{1−x}$P$_x$)$_2$ crystals was performed using a compact scanning electron microscope (SEM)—energy dispersive x-ray spectroscopy (EDS) device COXEM EM-30plus equipped with an Oxford Silicon-Drift-Detector (SDD) and AZtecLiveLite-software package. The EDS analyses on the BaNi$_2$(As$_{1−x}$P$_x$)$_2$ crystals revealed phosphorus content $x = 0.0, 0.035, 0.070, 0.075$, and 0.10 with a typical uncertainty of $\Delta x = \pm 0.05$.

Elastoresistance measurements

DC-elastoresistance measurements were performed following the method described in ref.$^{30}$ We used piezoelectrics from Piezomechanik GmbH (Part. No. Pst 150/5 x 5 x 7) and miniature strain gauges from Vishay Precision Group. The samples and strain gauges were glued to opposite faces of the piece using DevCon 5 mn DuPont 4929N or Hans Wolbring Leitsilber silver electrical resistance along the [100]tet direction using a Lake Shore 372 resistance bridge. For BaNi$_2$As$_2$, although we used an 8-contact geometry to extract the longitudinal and transverse elastoresistance during the same temperature cycle, each channel was measured one after the other, i.e., with current flowing along only one direction at a given time. As shown in Supplementary Fig. 4, this method gives very similar results compared to standard 4-contact geometry differential elastoresistance measurements. For P-substituted samples standard differential elastoresistance experiments were performed. Some of the freestanding resistance measurements were also done using a combination of Keithley 6221 current source and Keithley 2182A nanovoltmeter in delta mode. Either DuPont 4929N or Hans Wolbring Leitsilber silver paints were used.

Several strain sweeps were performed at each temperature to ensure the reproducibility of the extracted elastoresistance coefficients. A slow rate of 6 V/s was used to drive the piece. If possible cooling and warming elastoresistance measurements were recorded, without any significant difference. All the presented elastoresistance coefficients are extracted from linear-
in-strain fits. Second-order fits do not lead to any significant change in the extracted first-order coefficients.

DATA AVAILABILITY
All relevant data are available from the corresponding author upon reasonable request.

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AUTHOR CONTRIBUTIONS
M.F., M.L.T., and A.E.B. initiated the elastoresistance project. T.L. and A.-H.H. grew the single-crystal samples and performed the EDS analysis. M.F., P.W., and A.E.B. developed the experimental elastoresistance setup. M.F. performed the elastoresistance experiments and analyzed the data. C.M. realized the thermal expansion...
measurements. K.W. performed the specific heat measurements. T.L., S.M.S., and M.M. performed the X-ray diffraction experiments. M.F., C.M., and A.E.B. developed the interpretations. M.F. wrote the manuscript with input from all authors.

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