Temperature-dependent inter-plane resistivity, \( \rho_c(T) \), was measured for the iron-based superconductor BaFe\(_2\)(As\(_{1-x}\)P\(_x\))\(_2\) over a broad isoelectron phosphorus substitution range from \( x = 0 \) to \( x = 0.60 \), from non-superconducting parent compound to heavily overdoped superconducting composition with \( T_c \approx 10 \) K. The features due to structural and magnetic transitions are clearly resolved in \( \rho_c(T) \) of the underdoped crystals. A characteristic maximum in \( \rho_c(T) \), found in the parent BaFe\(_2\)As\(_2\) at around 200 K, moves rapidly with phosphorus substitution to high temperatures. At the optimal doping, the inter-plane resistivity shows \( T \)-linear temperature dependence without any cross-over anomalies, similar to the previously reported in-plane resistivity. This observation is in stark contrast with dissimilar temperature dependences found at optimal doping in electron-doped Ba(Fe\(_{1-x}\)Co\(_x\))\(_2\)As\(_2\). Our finding suggests that despite similar values of the resistivity and its anisotropy, the temperature dependent transport in the normal state is very different in electron and isoelectron doped compounds. Similar temperature dependence of both in-plane and inter-plane resistivities, in which the dominant contributions are coming from different parts of the Fermi surface, suggests that scattering is the same on the whole Fermi surface. Since magnetic fluctuations are expected to be much stronger on the quasi-nested sheets, this observation may point to the importance of the inter-orbital scattering between different sheets.
similar to previous observation of $T$-linear dependence in in-plane transport. This lack of significant features in either in-plane or inter-plane resistivity makes BaP122 system distinct from both electron-doped BaCo122 (linear $\rho_a(T)$ and cross-over $\rho_c(T)$) and hole doped BaK122, all with cross-overs in both $\rho_a(T)$ and $\rho_c(T)$. This difference suggests that three-dimensional character of the Fermi surface and normal state scattering are important in iron pnictides, and these are significantly different for different compounds and dopant species.

**EXPERIMENTAL**

Single crystals of BaP122 were grown from stoichiometric mixtures of Ba (flakes), and FeAs, Fe, P or FeP (powders) placed in an alumina crucible, sealed in evacuated quartz tube. It was heated to 1150-1200°C, kept for 12 hours and then cooled down slowly to 800°C at a rate 1.5°C/h. Platelet crystals had typically 0.3 to 0.7 mm$^2$ surface area, their $x$ value was determined using energy dispersive electron probe microanalysis (EDX).

Samples for the study were extensively characterized by polarized optics and magneto-optic techniques to look for possible inhomogeneity, as described in detail in Ref. 24. Inter-plane resistivity was measured using a two-probe technique, relying on the negligibly small contact resistance. The details of the measurement procedure were the same as in our previous studies on pure and transition metal doped Ba122 compounds, see Refs. 14, 15, 25, and 26 for details. In brief, samples typically had dimensions $0.5 \times 0.5 \times (0.02-0.1)$ mm$^3$ ($a \times b \times c$), all sample dimensions were measured with an accuracy of about 10%. The top and bottom $ab$-plane surfaces were covered with ultrapure Sn solder, as described in Ref. 24, forming a capacitor-like structure. Tin-soldering technique produced contact resistance typically in the 10 $\mu$Ω range. Four-probe scheme was used down to the sample to measure series-connected sample, $R_s$, and contact, $R_c$ resistance. Taking into account that $R_s \gg R_c$, contact resistance represents a minor correction of the order of 1 to 5%. This can be directly seen for our samples for temperatures below the superconducting $T_c$, where $R_s = 0$ and the measured resistance represents $R_c$. A tendency of the samples to cleave along $ab$ plane leads to a serious problem in inter-plane resistivity measurements. Even in visually perfect crystals, we frequently encounter partial cracks, leading to current redistribution in sample cross-section, and admixture of in-plane resistivity into measured inter-plane resistivity. To control this problem, we used as thin samples as were available and performed measurements of $\rho_c$ on at least 5 samples of each composition. In all cases we obtained qualitatively similar temperature dependencies of the electrical resistivity, as represented by the ratio of resistivities at room and low temperatures, $\rho_c(0)/\rho_c(300)$. The resistivity value, however, showed a notable scattering and at room temperature was typically in the range 1000 to 2000 $\mu$Ω cm, which is very similar to all transition metal doped Ba122, as well as for hole-doped BaK122.

**RESULTS**

Fig. 1 shows the main experimental result of this paper, a temperature-dependent inter-plane resistivity of BaP122 for several compositions from non-superconducting parent compound, $x=0$, through optimally doped, $x=0.33$ and $T_c=30$ K, to heavily overdoped, $x=0.60$ and $T_c=10$ K. For the sake of comparison the data are plotted on a normalized resistivity scale, $\rho_c(T)/\rho_c(300K)$, and offset downwards for increasing $x$.

![FIG. 1. (Color online) Temperature-dependent inter-plane resistivity of BaFe$_2$(As$_{1-x}$P$_x$)$_2$ for (top to bottom) $x=0$ (parent compound, black curve), underdoped compositions $x=0.23$ (green) and $x=0.25$ (blue), optimally doped $x=0.33$ (red) and overdoped $x=0.48$ (magenta) and $x=0.60$ (dark yellow). The data are plotted on the normalized resistivity scale, $\rho_c(T)/\rho_c(300K)$, and offset progressively downwards for higher $x$ to avoid overlapping. Arrows show a position of the resistivity cross-over temperature $T_{max}$ and of the structural, $T_S$, and magnetic $T_N$ transitions.](image-url)

Several features should be noticed. First, the curves for samples with $x=0.23$ and $x=0.25$ show a clear upward turn on cooling through the temperature of structural transition $T_S$ and downturn below the temperature of magnetic transition $T_N$, marked with arrows in Fig. 1. The values of $T_S$ and $T_N$ are in good agreement with NMR results. This splitting of structural and magnetic transitions in BaP122 is similar to electron-doped BaCo122.

An additional feature is clearly observed in $\rho_c(T)$ in parent and underdoped compositions $x=0.23$ and $x=0.25$.
at temperatures above 200 K. The $\rho_c(T)$ changes slope and shows a downturn on warming with resistivity taking very shallow maximum at a temperature $T_{\text{max}}$ as indicated with arrows. By comparison with NMR studies in BaCo122 [14], and with $\rho_c(T)$ for other transition metal substitutions [15], we previously assigned this maximum in transition metal doped Ba122 to the onset of carrier activation over the pseudogap. Similar assignment was suggested for the explanation of a maximum in $\rho_c(T)$ and a slope-saturation in $\rho_c(T)$ in hole-doped Ba122 and in NaFeAs [19][21]. Alternatively, the slope change in $\rho_c(T)$ of optimally doped BaK122 was explained in multi-band scenario [20].

In Fig. 2 we summarize a doping evolution of the characteristic temperatures of the c-axis resistivity: maximum $T_{\text{max}}$, temperatures of the structural, magnetic and superconducting transitions, for electron- (BaCo122) and isoelectron- doped (BaP122) BaFe$_2$As$_2$ compounds. For the latter we also show temperatures of nematic transition found in magnetic torque measurements [17]. The $\rho_c(T)$ maximum shows a dramatic asymmetry in $x$ for electron-doping and isoelectron substitutions. The crossover temperature is rapidly suppressed with doping in BaCo122, it is preceded by metallic temperature dependence at high temperatures above a minimum in $\rho_c(T)$ for heavily doped BaCo122. A close to $T$-linear $\rho_c(T)$ dependence is found at a critical concentration $x=0.313$ [14], and normal metallic $\rho_c(T)$, and temperature-independent Pauli susceptibility, $\chi(T)$, and Hall constant are restored for $x>0.313$ [14][40].

In stark contrast with both these doping dependences, crossover temperature $T_{\text{max}}$ shoots up with $x$ of isoelectron P-substitution, and this evolution leads to an interesting difference in the temperature-dependent anisotropic resistivity at optimal doping, as shown in Fig. 3. Two panels show $\rho_a(T)$ and $\rho_c(T)$ on a normalized resistivity scale, $\rho(T)/\rho(300K)$, for phosphorus isoelectron-substituted (top panel) and cobalt electron-doped (bottom panel) BaFe$_2$As$_2$. In both cases the resistivity above $T_c$ is close to $T$-linear, but the behavior at higher temperatures differs dramatically, and reveals a clear distinction: the crossover anomalies are absent in the isoelectron-substituted BaP122, while they affect only inter-plane transport in BaCo122.

**DISCUSSION**

In BaP122 both in-plane and inter-plane resistivities show the non-Fermi liquid $T$-linear dependence near the optimum doping $x \sim 0.3$, a concentration at which the antiferromagnetic quantum critical point has been observed [7]. At higher dopings, $\rho_c(T)$ becomes superlinear at low temperatures and increasing doping it gradually evolved towards the Fermi-liquid $T^2$ dependence, similar to the doping-evolution found for $\rho_{ab}(T)$ [5]. The $T$-linear resistivity near the QCP is consistent with the inelastic scattering by two-dimensional (2D) antiferromagnetic fluctuations [32], which are also evident from the Curie-like temperature dependence of the $1/(T_1T)$ (where $T_1$ is the NMR relaxation time) [7].

According to the band structure calculations, the Fermi surface of BaP122 comprises five sheets, three hole- and two electron [5][40]. These were observed experimentally by ARPES measurements [33][34] for the whole series of compounds. The warping, important for the inter-plane transport, is strongest for the hole- Fermi surface and it increases with $x$ in BaFe$_2$(As$_{1-x}$P$_x$)$_2$. This increased warping of the hole sheets has been observed by both ARPES studies [34] and quantum oscillations for phosphorus-rich compositions close to $x = 1$ [10]. The strongest curvature is found near the $Z$ point of the Brillouin zone, for the Fermi surface with dominant contribution of the $d_{z^2}$ orbital, which does not have significant nesting with the electron sheets and thus should be least affected by magnetic fluctuations. The in-plane conductivity is governed by the electron sheets with higher mo-
change the resistivity anisotropy beyond a factor of approximately two uncertainty of the geometric factors with phosphorus substitution from $x=0$ to $x=0.6$. Within this uncertainty, our resistivity anisotropy, $\gamma_p \approx 6\pm 2$, is in semi-quantitative agreement with the anisotropy of the upper critical field, $\gamma_H=1.44$ in ac magnetization measurements [37] and $\gamma_H=2.49$ as found in specific heat study [38], projecting to $\gamma_p = \gamma_H^2$ of about 2.1 and 6.2, respectively. The anisotropy of the upper critical field $\gamma_H$ shows mere 10% change with $x$ variation from 0.3 to 0.55 [38].

In our study of the inter-plane resistivity in transition metal doped Ba122 [14, 15] we observed a clear correlation of the maximum in $\rho_c(T)$ with onset of significant temperature dependence of the Knight shift in NMR measurements [12, 13]. Interestingly, this correlation extends to BaP122 as well, with the Knight shift being constant at the optimal doping [7], and no maximum being observed in $\rho_c(T)$, Fig. 1.

Infrared reflectivity measurements from the conducting $ab$-plane find pseudogap features at low temperatures in both optimally doped BaCo122 and BaP122 [16]. In both cases the feature appears on cooling in the temperature range between 100 and 200 K. While this temperature in BaCo122 is in reasonable agreement with the position of the $T_{\text{max}}$ crossover in $\rho_c(T)$, in BaP122 at optimal doping this maximum either moves to above 400 K or is suppressed completely. This fact may be suggestive that either the feature associated with the pseudogap in $ab$-plane spectroscopic reflectivity studies in BaP122 [16] is related with another anomaly in the normal state of iron pnictides, nematic response above $T_s$ [17, 18, 39, 40], or the pseudogap feature observed in the $ab$-plane reflectivity measurements in BaP122 does not affect transport along the $c$-axis. Further studies including Hall effect and optical conductivity measurements along the $c$-axis would be desirable to understand this point.

Comparison with the cuprates. The onset of the pseudogap feature in the cuprates is frequently determined from a temperature of deviations from $T_\text{linear}$, a dominant anomalous feature of the normal state transport closely linked with superconducting $T_c$ (see Ref. 42 for review). Similar analysis was recently done in BaP122 [17] and found a good coincidence with nematic feature found in torque measurements and inter-plane resistivity anisotropy in strain-detwinned samples [18]. In the cuprates the nematic order leads to a two-fold symmetry breaking in the plane as well [19, 41]. It was suggested that nematicity represents an order parameter for the pseudogap state in the cuprates [41].

CONCLUSIONS

Measurements of the inter-plane resistivity in BaP122 show that despite similar suppression of the mag-
netic/structural transitions with electron-doping and isoelectron-substitution into Ba122, the crossover maximum feature reveals dramatic difference in response between these cases. The presence/absence of the inter-plane resistivity maximum correlates with the presence/absence of the temperature-dependent NMR Knight shift in BaCo122/BaP122. Contrary to BaCo122, the inter-plane resistivity maximum in BaP122 shows no correlation to nematic anomalies of in-plane resistivity.

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