Point contacts are used to investigate the anisotropy of the superconducting energy gap in LuNi$_2$B$_2$C in the $ab$ plane and along the $c$ axis. It is shown that the experimental curves should be described assuming that the superconducting gap is nonuniformly distributed over the Fermi surface. The largest and the smallest gaps have been estimated by two-gap fitting models. It is found that the largest contribution to the point-contact conductivity in the $c$ direction is made by a smaller gap and, in the $ab$ plane by a larger gap. The deviation from the one-gap BCS model is pronounced in the temperature dependence of the gap in both directions. The temperature range, where the deviation occurs, is for the $c$ direction approximately 1.5 times more than in the $ab$ plane. The $\Gamma$ parameter, allowing quantitative estimates of the gap anisotropy by one-gap fitting, in the $c$ direction is also about 1.5 times greater than in the $ab$ plane. Since it is impossible to describe satisfactorily such gap distribution either by the one- or two-gap models, a continuous, dual-maxima model of gap distribution over the Fermi surface should be used to describe superconductivity in this material.

I. INTRODUCTION

The goal of this study was to investigate the anisotropy of the energy gap in nickel borocarbide superconductors ReNi$_2$B$_2$C. The crystallographic structure of these compounds resembles to some extent the structure of HTSC materials. It consists of alternating Re-$C$ and Ni$_2$-$B_2$ layers (Re is a rare earth metal). There is a point of view that the electron properties of ReNi$_2$B$_2$C compounds in the normal state are isotropic because of the strong carbon bond along the tetragonal $c$ axis. This is supported by the temperature dependence of isotropical resistivity in the single crystal of YNi$_2$B$_2$C. On the other hand, there is also data about a substantial anisotropy in these compounds. In the superconducting state the experimental data are also contradictory. The upper critical fields for $H$ parallel and perpendicular to the $c$ axis and the derived superconducting parameters do not show any anisotropy for the YNi$_2$B$_2$C single-crystal samples in agreement with magnetization and torque magnetometry measurements, but a small anisotropy is observed for the LuNi$_2$B$_2$C single crystals. Meanwhile, a very strong anisotropy of the superconducting energy gap was reported for LuNi$_2$B$_2$C in the $ab$ plane. This conclusion was based on the observation of delocalized quasiparticles in thermal conductivity at very low temperatures. According to this analysis, the smallest gap differs at least 10 times from the gap in other directions. Recently, evidence for the presence of nodes along (100) direction was provided by the field-angle thermal conductivity, field-angle heat capacity, and ultrasonic attenuation measurements of YNi$_2$B$_2$C.

In accordance with the experimental data an anisotropic superconducting gap function was proposed in Ref. 10 [see, also, Fig. 52 from the recent review (Ref. 11)]. In that model the gap has nodes along (100) directions and attains the maximum values along (110) directions. In the case of $s+g$ symmetry, the elastic scattering leads both to a decrease of $T_c$ and to an “isotropization” of the gap with vanishing nodes. We do not know any direct measurement of gap anisotropy involving the Andreev reflection. The STM tunneling measurements at 4 K report a gap of 2.2 meV along the $c$ axis, yielding a low low ratio $2\Delta/kT_c = 3.2$ for LuNi$_2$B$_2$C. By contrast, there are point-contact measurements of the gap in the $ab$ plane (e.g., Refs. 15 and 16), which yield $2\Delta/kT_c = 3.7–3.8$. The point-contact spectroscopy method in the latter measurements cannot, however, provide angular resolution of the gap anisotropy much better than $\pi/2$.

In the present investigation we have found that both in the $ab$ plane and in $c$ direction the experimental point-contact spectra cannot be fitted satisfactorily with the one-gap theoretical curve, even when broadened by an adjustable broadening parameter. Fitting the experimental spectra both at a small bias ($eV < \Delta$) and at a bias larger than $\Delta$ forces us to use at least the two-gap fitting curve like the one for the two-band superconductor. But even the two-gap fitting with the two proper broadening parameters was not good enough to approximate the experimental characteristic at the middle energy region ($eV < \Delta$). Moreover, for the two-band model one could expect transitions of Cooper pairs between the bands to introduce an additional depairing factor, which is analogous to other depairing factors like magnetic impurities, strong magnetic fields, etc. In such case the superconducting order parameter and the energy gap should differ from each other, which could be accounted for by using Ref. 18. Unfortunately even using the latter theory satisfactory fits could not be obtained. Therefore we suggest that only a continuous gap distribution with the two energy gap maxima around the Fermi surface could satisfactorily approximate the experimental spectra at low temperature ($T \ll T_c$). We have determined tentatively the value of these maxima and the range of gap distribution which corresponds well to the recent STM measurements.
II. EXPERIMENTAL TECHNIQUE

The point contact measurement was performed on single crystal LuNi$_2$B$_2$C with $T_c = 16.9$ K grown by Canfield and Bud’ko using a flux method. Geometrically, our crystal was a thin (0.1 – 0.2 mm) plate with the c axis perpendicular to its plane. The single crystal surface always contains quite a thick layer in which superconductivity is either absent or strongly suppressed. To perform measurement in the ab plane, the crystal is usually cleaved and the point contact is made between a metallic counter electrode and the cleaved surface. It was technically problematic to produce a cleavage perpendicular to the c direction. In this case the crystal surface was cleaned with a 10% HNO$_3$ solution in ethanol. As the measurement in the ab plane shows, both cleavage and etching yield identical results.

The point contact in the ab plane is fabricated between an edge of the silver electrode and a freshly cleaved (or etched) corresponding facet of the single crystal. The deviation from the perpendicular to the c direction might amount to about $5^\circ$ – $10^\circ$. We do not know a priori along which of the in-plane directions the contact is obtained. But since we used a selection rule to choose the highest observable superconducting energy gap with the largest nonlinearity at the gap double-minimum structure, the contact axis is presumably along the directions, where the in-plane gap is maximum. To produce contacts in the c direction the traditional “needle-anvil” geometry is used. The radius of the needle is about 1 – 3 micron. The temperature was measured using a special cryogenic insert (its close analogue is described elsewhere).

The point contact resistances varied typically from several Ohms to tens of Ohms. For detailed investigations we chose the point contacts with the highest permissible tunneling which was controlled against the differential resistance maximum at zero bias and by the maximum nonlinearity in the dV/dI-double-minimum region, corresponding to the intact superconducting surface under the contact. Unfortunately, a complete set of curves in the whole $T_{min}$ – $T_c$ interval was obtained only for a few contacts. Because of their rather high resistance and the long-duration (over 10–12 hours) of the measurements, many of the contacts were broken down. The temperature interval ranges from the minimal available $T$ (~1.5 K) to $T_c$.

III. RESULTS AND DISCUSSION

Some curves of the first derivatives dV/dI measured at different temperatures on LuNi$_2$B$_2$C-Ag point contacts along c axis and in the ab plane are shown in Figs. 1 and 2.

The measured curves were symmetrized $dV/dI_{sym} = 1/2[dV/dI(V) + dV/dI(-V)]$ and normalized to the normal state at $T > T_c$, except in Figs. 1 and 2, where the raw data are shown. Each curve contains several hundred of experimental dots and at the scale shown in the presented figures, all the curves have negligible noise. In other figures experimental curves are shown by dots whose number is decreased in order to be discerned. Statistically (several tens of contacts were examined for each direction), the distance between the minima in the first derivative $dV/dI$ characterizing the average value of the gap was 12–15% larger in the ab plane compared to that in the c direction. This is a very crude estimation of the $c \rightarrow ab$ anisotropy.

The theoretical predictions and the experimental results were compared using two approaches. First, a model was applied, which describes electrical conductivity of pure $S$-c-N point contacts in the presence of an arbitrarily transparent potential barrier at the boundary between the metals. This model allows for the finite lifetime of the Cooper pairs. The I-V characteristics of the point contact are described as follows:

$$I = \frac{1}{2eR_N} \int_0^\infty d\varepsilon \left[ \tanh \frac{\varepsilon + eV}{2T} - \tanh \frac{\varepsilon - eV}{2T} \right] \times \frac{1}{(2d^{-1} - 1) + g_{ee}[1 + g_{ee}^2 + |f|^2 + 4(d^{-1} - 1)Re g_j]} ,$$

where $g_{ee} = u/\sqrt{u^2 - 1}$, $f = 1/\sqrt{u^2 - 1}$, $d = 1/(1 + Z^2)$, $d$ is the transparency of the potential barrier, and $Z$ is the tunneling parameter. The value $u = \varepsilon/\Delta_c$ ($\Delta_c$ is a complex energy-independent order parameter of the superconductor), which can be found from the following equation:

$$u = \frac{\varepsilon}{\Delta} + i\frac{\gamma}{\sqrt{u^2 - 1}} ,$$

where $\Delta$ is the BCS order parameter obtained from

FIG. 1. Differential resistance of LuNi$_2$B$_2$C-Ag point contact at different temperatures. The contact axis is parallel to c. Not to overload the figure, we give only several representative curves.
and $\gamma = 1/\tau_5 \Delta$ is the pair breaking parameter ($\tau_5$ is the mean free time during, e.g., spin-flip scattering at impurities). When magnetic impurities are absent, $\tau_5$ tends to infinity and Eq. (1) coincides with the equation for current in Ref. 24. The energy gap $\Delta_0$ and the order parameter $\Delta$ are related as follows:

$$\Delta_0 = \Delta (1 - \gamma^{(3)/2})^3.$$

The other approach was based on the generalized Blondin-Tinkham-Klapwijk (BTK) model, commonly used to describe $S-c-N$ point contacts. The model allows for the finite lifetime of quasiparticles $\tau = h/\Gamma$ determined by inelastic scattering, which leads to the broadening of the density of states in the superconductor. Formally, according to the theory the BTK-based results are obtained under the condition of strong pair breaking ($|\alpha| \gg 1$). Therefore, in the strict sense, the generalized BTK model contains no gap. For any infinitesimal broadening parameter $\Gamma$, at $T = 0$ the density of states near the Fermi surface is nonzero. In theory, the order parameter is a quantity analogous to the pseudogap in the generalized BTK model. In the following we use terms “order parameter” and $\gamma$ for the model of Ref. 18, while the “gap” and $\Gamma$ for the commonly used BTK model, except special cases, where we refer to $\Delta_0$ related to the model.

Now we consider more closely the methods of fitting the theoretical curves to the experimental results. The iteration method commonly used in this case is quite good for small broadening. However, when the broadening $\Gamma$ is comparable with the gap ($\gamma > 0.3$), the results thus obtained are ambiguous and often dependent on the starting $\Delta, \Gamma$ (or $\gamma$) and $Z$ values. In our calculation, we therefore used the technique of coordinate descent with a postponed solution.

First, we specified an interval in which $\Delta$ is searched at a given temperature. The interval was then subdivided into equidistant parts, $\Delta_1, \Delta_2, \ldots, \Delta_n$, and $\gamma$ and $Z$ were fitted for each $\Delta_i$. The procedure was as follows: after each step of $\gamma$-fitting, a complete $Z$-fitting was performed and then the next step of $\gamma$-fitting was considered. Before each calculation of the average rms deviation $F(\Delta)$, the amplitudes of the fitting and the experimental curves were made equal by multiplying the $\gamma$-coordinate of the fitting curve by a scale-factor $S$ and then shifting it along the $\gamma$-axis by an amount $B$. The values of $B$ and $S$ were chosen to minimize the deviation $F(\Delta_i)$. The standard algorithm for determination of $B$ and $S$ is considered, for instance, in Ref. 26. As a result, for each $\Delta_i$ we found $\gamma_i$ and $Z_i$, at which the difference between the shapes of the theoretical and experimental curves characterized by the rms deviation $F(\Delta_i)$ was the smallest one. The same holds for $\Gamma$ in the BTK fitting model.

The calculation for some temperatures is shown in Fig. 3. It is evident that at $T_c$ the technique cannot ensure unambiguous results. At 14.5 K, the curve $F(\Delta)$ is practically horizontal. The determination of the $\Delta$-value should take into account other factors as well. For example, it is important that the tunneling parameter $Z$ and the scale factor $S$ should be invariant. The dependences $S(\Delta)$ and $F(\Delta)$ at $T = 7.5$ K are shown in Fig. 4. It is seen that the minimum $F$ corresponds to a certain $S$ value. The BTK calculation of the corresponding dependences at the same temperature yields broader $F(\Delta)$ curves. Nevertheless, the minima positions in these curves are quite definite.

At low temperatures the shapes of the theoretical and the experimental curves were in rather poor agreement for all of the borocarbides investigated (Re=Er, Dy, Tm, Lu) irrespective of the point contact orientation. The temperature dependence of the rms deviation in shape in the $c$ direction is shown in Fig. 5 for both models. It is seen that the best agreement is achieved above 7 K. At $T < 7$ K, the largest discrepancy between the shapes is observed in the region of extrema. The central maximum of the differential resistance in the experimental curve is considerably narrower than that in the best fit, whereas the minima are shifted to higher bias (Fig. 6).

Some experimental curves have a kink marked with an arrow in Fig. 6. This shape is typical of a superconductor with two gaps of close energies. The temperature dependence of the broadening (pair breaking) parameter also supports the existence of two gaps. Since Lu is a nonmagnetic material, it is natural to expect a negligible broadening (pair breaking) parameter for the Lu-based borocarbide point contacts. However, that is not the case, as is evident from the temperature dependences (Fig. 7) of the broadening (pair breaking) parameters obtained within the two models (Ref. 18 and BTK).

In our calculation, we therefore used the technique of coordinate descent with a postponed solution.
Note that $\gamma$ and $\Gamma$ are different quantities. In the generalized BTK model\textsuperscript{25} $\Gamma$ is independent of the gap and has the gap dimensions; whereas, in the model of Ref. 18, $\gamma$ is described as $\gamma=1/\tau_0$, and, for $\gamma<1$, $\gamma$ is related to the gap $\Delta_0$ and the order parameter $\Delta$ as $\gamma=[1-(\Delta_0/\Delta)^{2/3}]^{1/2}$. As the temperature is increased, both $\Gamma$ and $\gamma$ decrease, and, at temperatures $>7.5$ K for the $ab$-plane and $>11$ K for the $c$-direction, tend to zero. Meanwhile, the one-gap fit starts to approximate the experimental curve more and more accurately. In this case the one gap fit with the broadening (pair breaking) parameter is a tool of describing a certain average gap (order parameter).

The largest and smallest gaps can be estimated by fitting different portions of the experimental curve. In this case, to match the amplitudes of the fitting and the experimental curves, we used another method. Namely, the $\gamma$-coordinates at the central maximum and near the minima are scaled to coincide (Fig. 8). The best agreement in the central maximum region can give us an estimate of the smallest gap (curve 1), while the best agreement in the “wing” region yields the largest gap (curve 2). By “wings” we imply the portion of the experimental curve at biases higher than the energy gap double-minimum structure in the $dl/dI$ characteristic, namely at $eV \approx 3$ meV. The estimates however are rather rough.

Further, we follow the method of approximation of $I$-$V$ characteristics, which was calculated for the $S$-$I$-$N$ contacts of MgB$_2$ within a two-band model in Ref. 17. There, the total conductivity of the tunnel contact is a sum of the $\sigma$- and $\alpha$- band conductivities analyzed by applying the BTK model. To describe the resulting curve, we also used a model of two independent parallel-connected point contacts with different gaps whose conductivities are additive. The contributions of these conductivities account for the part of the Fermi surface containing a particular gap. Thus, for the two-gap model an experimental curve is fitted by the following expression:

$$\frac{dV}{dl} = \frac{S}{dV/d\Delta_1, \gamma_1, Z} + \frac{dV/d\Delta_2, \gamma_2, Z(1-K)}{1-K}.$$ 

With a proper choice of the coefficient $B$, here the coefficient $K$ reflects the contribution of the part of the Fermi surface having the gap $\Delta_1$, $S$ is the scaling factor discussed for the one-gap approximation. To obtain the best agreement with the experiment, the parameters used in this expression are allowed to differ from those found for fitting of the separate portions of the experimental curve. It appears that when...
TABLE I. The fitting results for superconducting parameters along two mutually perpendicular directions, using Ref. 18 and the BTK theory (Ref. 25). $\Delta_1$, $\Delta_2$, are the order parameters in the two-gap fit; whereas, the subindex 0 corresponds to the energy gaps in the model of Ref. 18. The pair breaking parameters $\gamma_{1,2}$ and the broadening parameter $\Gamma_{1,2}$ correspond to Refs. 18 and 25, respectively. $Z$ is the tunneling parameter. The temperature corresponds to that of the experiment. The relative contributions of each superconducting order parameter and gap are given in percent. $\Delta$ and $\Gamma$ are given in meV. $\gamma$ and $Z$ are dimensionless.

<table>
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<td>1.48</td>
<td>1.47</td>
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the contribution of the smaller gap prevails, the “wings” fit gives smaller values. In this case, the best result is achieved with the two-gap fit (Fig. 8, curve 4) using the parameters for curves 1 and 3, the latter is marked with dots. Correspondingly, if the larger gap prevails, the central-maximum fit gives higher values. In Fig. 6 (dotted curve) an example of the two-gap fit for the $ab$-plane is shown.

Although the two-gap fit shows much better agreement in the regions both of the central maximum and the “wings” (see Figs. 6 and 8), it cannot provide a complete description of the experimental curves, especially at the double-minima structure of $dV/dI$, and in the $c$ direction (Fig. 8). Most likely, this is because of a continuous distribution of the gaps over the Fermi surface.

Our estimation of point-contact parameters in the $ab$ plane and in the $c$ direction was made within one-gap and two-gap models using the equations of Ref. 18 and BTK. The results are presented in Table I. $\Delta_1$, $\Delta_2$, are the highest and lowest order parameters (the same as BTK gaps) from the two-gap fit; $\Delta_01$ and $\Delta_02$ are the energy gaps corresponding to $\Delta_1$, $\Delta_2$; $\gamma_1$, $\gamma_2$ are the pair breaking parameters of the model for the largest and smallest gaps. The same holds for the widening parameters $\Gamma_1$, $\Gamma_2$ in the BTK model. Independently determined $Z$’s are the tunneling parameters, which are the same for the largest and smallest gaps and are thus quite self-consistent. In Table I the order parameters (gaps) based on the two-gap fit appear along with the contributions to the total conductivity from the Fermi surface region with the corresponding gap. For example, for the $c$ direction contact, the contribution to the conductivity is 40% from the region with an order parameter of 3.2 mV and 60% from the region with 1.82 mV.

It is seen that within both the models, the dominant contribution to the total conductivity in the $c$ direction is made by the Fermi surface region where the order parameter is lower. The higher order parameter makes the dominant contribution in the $ab$ plane. On the average this correlates with gaps in $ab$ and $c$ directions predicted by the $(s+g)$ model.

Although the part-by-part fitting is rather rough, we tried this procedure for tracing the temperature dependences of the smallest and largest gaps. However, more or less definite values were obtained only at the lowest temperatures because the smallest gap is estimated within a relatively small part of the experimental curve (near zero bias). As for the largest gap, its temperature dependence was traced up to the moment when the calculations over the entire curve and over the “wings” start to give similar results.

The temperature dependences of the scale factor $S$ in the $c$ direction are shown in Fig. 9. They were found for the largest gap (“wing” fitting is marked with “max” throughout) and for the average gap (entire-curve fitting) by the equations of Ref. 18 and BTK. The temperature dependences of the tunneling parameter $Z$ are shown for both models. To avoid crowding in the figure, only the $Z$-values obtained from the

FIG. 9. Temperature dependences of the tunneling parameter $Z$ and the scale factor $S$ for the LuNi$_2$B$_2$C-Ag point contact in the $c$ direction obtained with models from Refs. 18 and 25. The scale factor was found from fitting over the entire curve and over the “wings” (scaling max).
FIG. 10. Temperature dependence of the order parameter $\Delta$ and energy gap $\Delta_0$ for the one-gap fitting models (Refs. 18 and 25) in the $c$ direction. The $T$-dependence of the largest order parameter, indicated by max, was estimated from fitting over the “wings.”

The calculation was made both over the entire curve and the “wing” one-gap fittings, 18,25 are shown in Fig. 10 along with the BCS curve. As in the case of the scaling factor $S$ (Fig. 9), deviation starts at $T \leq 11$ K, while at higher temperature all the fitting procedures result in the same BCS-like dependence.

The temperature dependences of the order parameter and the gap in the $c$ direction, obtained both by the entire-curve and the “wing” one-gap fittings, 18,25 are shown in Fig. 10 along with the BCS curve. As in the case of the scaling factor $S$ (Fig. 9), deviation starts at $T \leq 11$ K, while at higher temperature all the fitting procedures result in the same BCS-like dependence.

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This procedure includes manually offsetting dots of the $K$ fitting curve in the intermediate steps of the adjustment. The broadening parameter $s$ equals $D$, the Fermi surface with the gap $K$ where the discrepancy is the highest. Its contributions, where the order parameter coincides with the energy gap. As a first approximation, it is supposed that the coefficients $Z$ are the same, and the order parameter coincides with the energy gap.

Let us illustrate this approach by fitting the experimental curves for the $c$- and $ab$-directions, respectively. One can see that the multigap fit with eight fitting parameters and a continuous distribution of the gaps matches the experimental curve very well. One should not trust this fitting literally, but it shows that the width of the gap distribution can be quite appreciable, down to small values, which agrees with the tunneling results and theoretical $(s+g)$ model of the gap nodes in [100] directions.

**IV. CONCLUSION**

In this study the anisotropy of the superconducting energy gap was measured for the first time on LuNi$_{2}$B$_{2}$C in the $ab$ plane and $c$ directions. It is found that at low temperatures the experimental curves should be described assuming a gap distribution over the Fermi surface. Within the two-gap model, the largest contribution to the total conductivity is made by the Fermi region with a smaller gap in the $c$ direction and by the Fermi region with a greater gap in the $ab$ plane. At $T = 1.5$ K the largest and the smallest order parameters in the $c$ direction are $\Delta_{\text{max}} = 3.2$ meV, $\Delta_{\text{min}} = 1.82$ meV; in the $ab$ plane these are $\Delta_{\text{max}} = 3.03$ meV, $\Delta_{\text{min}} = 2.0$ meV. An attempt to fit the low temperature experimental curves connected point contacts having different gaps. Their contribution to the total conductivity can qualitatively account for the region of the Fermi surface containing a particular gap, as was considered in Ref. 27.

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**FIG. 12.** Modeling of the anisotropic superconducting energy gap distribution over the Fermi surface with multiple parallel junctions of different gaps. The contribution of a particular energy gap correlates with its part over the Fermi surface. The distribution of a particular contribution to the total conductivity is described by the $K_i(\Delta_i)$ dependence. The shape of the distribution $K_i(\Delta_i)$ is simulated by superposition of two peaks. Each peak is described by the following parameters: the energy of the central maximum $\Delta_i$ or $\Delta_{p_i}$, half-width of the right and left slopes $\sigma_1$, $\sigma_2$ or $\sigma_5$, $\sigma_4$, and the heights $h_1$ and $h_2$, correspondingly. By variation of these eight parameters we attempt to reach the best coincidence between the experimental and theoretical curves.

![Graph](image)

**FIG. 13.** Approximation of the experimental curve by the use of the multigap model for the $c$ direction, $T = 1.47$ K, $Z = 0.559$. Inset, the contribution to the total conductance from different parts of the Fermi surface with different gaps. The distribution is modeled by 31 equidistant points, $\Delta_{\text{min}} = 0.7$ meV, $\Delta_{\text{max}} = 4$ meV. The resulting curve exhibits two peaks with maxima at 1.8–2 meV and 2.5 meV.

![Graph](image)

**FIG. 14.** Approximation of the experimental curve by the use of the multigap model for the $ab$ plane, $T = 1.48$ K, $Z = 0.755$. Inset, the contribution to the total conductance from different parts of the Fermi surface with different gaps. The distribution is modeled by 31 equidistant points, $\Delta_{\text{min}} = 1.0$ meV, $\Delta_{\text{max}} = 3.35$ meV. The resulting curve exhibits two peaks with maxima at 2 meV and 3.1 meV.
shows that the gap is distributed starting from 0.8 meV. This corresponds well to the recent STM observations in Ref. 19 discussed above. We have found the deviation of the temperature dependent gap from the BCS theory for both tested directions. This deviation is connected with the impossibility to describe the gap distribution in terms of the one-gap model, at least at low temperatures. The temperature range where the described deviation is observed for the $c$ direction, is about 1.5 times greater (1.5–11.5 K) than that in the $ab$ plane (1.5–7.5 K). The broadening parameter $\Gamma$, allowing quantitative estimation of the degree of anisotropy, is also bigger in the $c$ direction.

The experimental results are described on the basis of the generalized BTK model\textsuperscript{25} and the Beloborod’ko theory,\textsuperscript{18} considering the electrical conductivity of ballistic $S$-$c$-$N$ point contacts in the presence of an arbitrarily penetrable potential barrier and allowing for the finite lifetime of Cooper pairs. For superconductors with a multiband electronic structure, the interband transitions of Cooper pairs may lead to their finite lifetime. Previously, a two-band model was suggested for nickel borocarbide superconductors.\textsuperscript{28} The theory,\textsuperscript{18} which accounts more accurately for the force of pair breaking, may explain the difference between the order parameter and the gap.

Our next publication will concern the compound $\text{ErNi}_2\text{B}_2\text{C}$ which exhibits a magnetic transition near $T = 6$ K. In this context a detailed analysis and comparison of the two theoretical approaches\textsuperscript{18,25} are of paramount importance for understanding the results that can be obtained on magnetic $\text{ErNi}_2\text{B}_2\text{C}$.

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29 “fit” stands for the theoretical $dV/dI(V)$-dependence fitted to the experimental results.