Rutile Mineral Chemistry and Zr-in-Rutile Thermometry in Provenance Study of Albian (Uppermost Lower Cretaceous) Terrigenous Quartz Sands and Sandstones in Southern Extra-Carpathian Poland

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Abstract: The geochemistry of detrital rutile grains, which are extremely resistant to weathering, was used in a provenance study of the transgressive Albian quartz sands in the southern part of extra-Carpathian Poland. Rutile grains were sampled from eight outcrops and four boreholes located on the Miechów, Szydłowiec, and Pulawy Segments. The crystallization temperatures of the rutile grains, calculated using a Zr-in-rutile geothermometer, allowed for the division of the study area into three parts: western, central, and eastern. The western group of samples, located in the Miechów Segment, is characterized by a polymodal distribution of rutile crystallization temperatures (700–800 °C; 550–600 °C, and c. 900 °C) with a significant predominance of high-temperature forms, and with a clear prevalence of metapelitic over metamafic rutile. The eastern group of samples, corresponding to the Lublin Area, is monomodal and their crystallization temperatures peak at 550–600 °C. The contents of metapelitic to metamafic rutile in the study area are comparable. The central group of rutile samples with bimodal distribution (550–600 °C and 850–950 °C) most likely represents a mixing zone, with a visible influence from the western and, to a lesser extent, the eastern group. The most probable source area for the western and the central groups seems to be granulite and high-temperature eclogite facies rocks from the Bohemian Massif. The most probable source area for the eastern group of rutiles seems to be amphibolites and low temperature eclogite facies rocks, probably derived from the southern part of the Baltic Shield.

Keywords: heavy mineral analysis; trace elements; Lower Cretaceous; provenance; mature sediment; longshore current; Bohemian Massif; Baltic Shield

1. Introduction

The uppermost Early Cretaceous (Middle and Late Albian, c. 110.8–100.5 Ma) and Late Cretaceous epicontinental basin in Poland, called the Polish Basin, was part of the vast Central European Basin System (CEBS, Figure 1) [1]. The CEBS developed after the Variscan orogeny (c. 300 Ma) and extended from the North Sea to Poland [2]. The processes that took place in the CEBS from the Permian to the Palaeogene were successive marine transgressions and regressions with the dominant characteristics of shallow shelf sea sedimentation. In the late Jurassic, a significant uplift of Precambrian and Variscan structures was reported, e.g., [2,3] including SW Poland. After the Neo-Cimmerian tectonic phase, during the Earliest Cretaceous, the sea-level was low [4] and large land areas of extra-Alpine Europe had been elevated [5]. Sedimentation in the Polish Basin was restricted to the narrow Mid-Polish Trough, including terrestrial and shallow shelf facies. Additionally, the epicontinental Polish basin was isolated from the Tethys Sea during most of the Early Cretaceous [6–8]. In the Middle and Late Albian (latest Early Cretaceous) in the Polish Basin, as in everywhere in Europe, a significant eustatic marine transgression
began [9–11]. The sea transgressed from the west and merged with the Russian sea in the east [9]. The detrital material came, most probably, from nearby uplifted areas. The source areas (Figure 1) could have been (i) the Bohemian Massif S and SW of the study area; (ii) the Ukrainian Shield [5,9,12]; (iii) a hypothetical Holy Cross Mountain—Dobruja Land (HCMDL) [13], also called the Krukienic Island [14]; or (iv) the crystalline basement of the southern part of the Baltic Shield. Quartz as a main component of the Albian sands has little diagnostic significance in determining its provenance. Therefore, the analysis of heavy minerals seems to be a promising approach in provenance studies of these mineralogically mature sediments [15–19].

Figure 1. Albian palaeogeography of epicontinental sea and adjacent western Tethys of Europe with simplified distribution of facies (modified from [5,20]); 1—land areas; 2—deltaic, coastal, and shallow marine clastic facies (sands and conglomerates, sands and shales); 3—shallow-marine facies (sands, marls, carbonate marls, marly carbonates, carbonates); 4—mainly shallow-carbonate marine facies (marly carbonates, carbonates, chalk, white chalk, carbonate shales); 5—active fold belts, high relief; 6—marine carbonates; 7—deeper marine clastics (sand and shales); 8—deeper marine carbonates (with sands and shales); 9—carbonates and sands; 10—deeper marine facies (sands and shales, shales); 11—deeper marine shale facies from a rift area; 12—area of the study; HCMDL—hypothetic Holy Cross Mountains—Dobruja Land; ?—no precise data for the southern edge of the HCMDL.

Recently, the majority of provenance studies of sedimentary rocks containing clastic material have been based on the composition of heavy minerals [15,21,22]. The development of EPMA techniques has made it possible to carry out relatively inexpensive, rapid, and widely accessible analyses of trace elements in heavy minerals. This has con-
tributed to the use of heavy minerals in provenance analyses of siliciclastic deposits and the
determination of the host rocks and source areas in many places around the world [23–25].

Rutile, apart from zircon and tourmaline, is classified as a heavy mineral extremely
resistant to weathering [24] and is very common in clastic sediments, including quartz
arenite. Despite this, rutile is rarely used in provenance analyses of clastic material. It
is, however, an especially useful mineral in pure quartz arenites, which are characterized
by high ZTR (zircon-tourmaline-rutile) index, low quantity, and low diversity of heavy
minerals [18,26,27]. However, it is interesting because rutile composition can be used as a tool
in provenance studies [28–30]. The advancement of analytical techniques led to the recognition
of a relationship between the crystallization temperature and Zr content in rutile [31,32]. This
in turn allowed the calibration of the Zr-in-rutile (ZIR) geothermometer, which can also be
quite widely used in determining the provenance of detrital rutile [28,29,31,33,34].

The aims of this paper are as follows: (1) to comment on the physical properties
of rutile as an important factor influencing the mode of preservation of rutile in detrital
settings; (2) to report on the chemical composition of rutile from twelve outcrops and/or
boreholes scattered in extra-Carpathian Albian (uppermost Lower Cretaceous) sands of
southern Poland; (3) to test an application of the Zr-in-rutile thermometry in provenance
studies of quartz arenite; and, finally, (4) to point out some of the possible provenances of
rutile in the Albian quartz arenites to determine the host rocks and location of source area(s).

2. Regional Geological Setting

The study area covers the southern part of extra-Carpathian Poland where Albian
deposits are available partly as outcrops and partly in boreholes (Figure 2). The Albian
deposits crop out at the marginal parts of the Miechów Segment of the Szczecin-Łódź-
Miechów Synclinorium (Figure 2, Table 1). In the south-western part of the Miechów
Segment, they form a narrow, nearly 100 km long longitudinal belt of outcrops, including
the Korzkiew, Glanów-Stronczki, Przychody, Lełów, and Mokrzesz localities studied here.
In the north-eastern part of the Miechów Segment, close to the Holy Cross Mountains
(HCM), the Albian deposits also form a narrow, longitudinal belt, but with rather occasional
outcrops [35]. The best known outcrops in this part of the Miechów Segment occur at
Wrzosówka Hill (292.6 m a.s.l.) near Bolmin (studied here) and in an overburden active
quarry in Małogoszcz (approx. 5 km west of Bolmin). The Radomsko Folds with the
Chełmo Mount (in Polish: Góra Chełmo, 323 m a.s.l.) terminate the Miechów Segment from
the north and separate it from the Mogilno-Łódź Segment (Figure 2). One more outcrop
studied here, called Kopiec, is situated on the northern edge of the Annopol Anticline in
the Szydłowiec Segment of the southern part of the Mid-Poland Anticlinorium (Figure 2,
Table 1). The other four samples studied here come from boreholes in eastern Poland. The
thickness of the Albian sands in the Potok IG-1 borehole in the Szydłowiec Segment in the
southern part of the Mid-Polish Anticlinorium is approximately 20 m [36] and was sampled
from a depth of 248.0–249.7 m. The other three sampled boreholes, Tomaszów Lubelski
IG-1, Piaski IG-2, and Łuków IG-1, are situated in the vast tectonic unit called the Puławy
Segment in the southern part of the Kościerzyna-Puławy Synclinorium (Figure 2, Table 1).
The Tomaszów Lubelski IG-1 borehole is very close to the Mid-Polish Anticlinorium. The
thickness of Albian sands in this borehole is c. 12 m [37] and the core material was sampled
at the depth of 1006.7–1013.0 m. The thickness of Albian sands in the Piaski IG-2 borehole
is only c. 2.5 m [38] and the core was sampled at the depth 725.4–727.4 m. The Łuków
IG-1 borehole is located close to the Mazury-Podlasie Homocline (Figure 2, Table 1). The
thickness of Albian sandstone in the northernmost borehole, Łuków IG-1, is c. 25 m [39]
and was sampled at a depth of 510.1–524.6 m.
The Albian deposits directly cover Upper Jurassic limestones or calcareous clays at eleven locations [9,41]. Only in one borehole, Potok IG-1, are the calcareous sands overlying the pre-Albian Lower Cretaceous sand, sandstones, and calcareous sandstones [36]. In all cases, excluding the Chelmó Mount, the Albian sands and sandstones are covered by lithified carbonate (mainly inoceramid limestone) Cenomanian deposits [35,42]. At Chelmó Mount, the Albian sandstone is covered by Cenomanian glauconitic sands [43]. The thickness of the Albian deposits ranges from 0 to c. 50 m [42] in the south-western margin of the Miechów Segment, and approximately 26–76 m in the north-eastern margin [44]. The thickness of Albian sands and sandstones at Chelmó Mount [43], recorded as 180 m, seems to be overestimated. Previous investigations [6] show a maximum thickness of 100 m for the Albian at this locality. The thickness of Albian deposits in the Kopiec outcrop is 6.5 m and, in boreholes, as stated above, it ranges from c. 2.5 m (Piaski IG-2) to c. 25 m (Łuków IG-1). The differences in thickness of the Albian deposits were probably controlled
by active synsedimentary block faulting and a diversified pre-Albian surface morphology, which played a substantial role in controlling the local deposition of clastic material [35,42].

| Sample | Type of Locality | Locality            | Latitude     | Longitude   |
|--------|------------------|---------------------|--------------|-------------|
| KRZ    | outcrop          | Korzkiew            | 50°09′42.5″ N | 19°52′46.7″ E |
| GLA    | outcrop          | Glanów-Stroniczki   | 50°18′59.9″ N | 19°47′32.7″ E |
| PRZ    | outcrop          | Przychody           | 50°30′22.0″ N | 19°43′42.6″ E |
| LEL    | outcrop          | Lelów               | 50°41′48.4″ N | 19°36′18.9″ E |
| MO     | outcrop          | Mokrzesz            | 50°48′09.5″ N | 19°23′34.4″ E |
| GCH    | outcrop          | Chelmo Mount        | 51°03′27.2″ N | 19°44′38.3″ E |
| BOL    | outcrop          | Bolmin              | 50°48′36.5″ N | 20°20′37.1″ E |
| PO     | borehole         | Potok IG-1          | 50°45′43.1″ N | 22°19′13.1″ E |
| KOP    | outcrop          | Kopiec (Annopol)    | 50°54′27.0″ N | 21°50′28.1″ E |
| TL     | borehole         | Tomaszów Lubelski IG-1 | 50°24′59.8″ N | 23°34′38.6″ E |
| PK     | borehole         | Piaski IG-2         | 51°09′25.3″ N | 22°55′03.6″ E |
| LK     | borehole         | Łuków IG-1          | 51°56′32.9″ N | 22°34′33.7″ E |

### Table 1. GPS coordinates of the examined localities.

### 3. Materials and Methods

Twelve localities within the Albian deposits were sampled in the extra-Carpathian area of southern Poland (Table 1, Figure 2). Sand and sandstone samples were taken from seven natural outcrops: Korzkiew (KRZ), Glanów-Stroniczki (GLA), Przychody (PRZ), Lelów (LEL), Mokrzesz (MO), Bolmin (BOL), and Chelmo Mount (GCH) of the Miechów Segment. A sand sample from Kopiec (KOP) near Annopol and one sand sample from the Potok IG-1 borehole (PO) from the Szydłowiec Segment were also taken. Two sandy samples from the Tomaszów Lubelski IG-1 (TL) and the Piaski IG-2 (PK) borehole and one quartz sandstone sample from the Łuków IG-1 (LK) borehole, all from the Pulawy Segment, were also taken.

In order to separate the heavy minerals, a 1 kg sample from each outcrop (KRZ, GLA, PRZ, LEL, MO, GCH, BOL, KOP) and 0.5 kg sample from each core material (TL, PO, PK, LK)—the weight of the core sample was limited owing to the limited amount of material available for sampling—were used. The sandstone samples from the Chelmo Mount were initially crushed in a Testchem LKS-60 jaw crushe. All twelve samples (loose sands and crushed material) were then rinsed with double-distilled water to dispose of the clay fractions, three magnetic fractions (0.3 A, 0.8 A, 1.2 A) and then sieved to obtain 63–125 µm size fractions. From these fractions, three magnetic fractions (0.3 A, 0.8 A, 1.2 A, and a diamagnetic one at 1.2 A) were obtained from a single sample after separation on the Franz LB-1 Magnetic Barrier Laboratory Separator. The Frantz isodynamic separator was set to standard operating conditions, with the side slope fixed at 15° and the chute forward slope set to 20° [45]. The most magnetic minerals, such as Fe-Ti oxides and some garnets, tend to concentrate in the 0.3 A fraction, while rutile falls into the diamagnetic fraction, and only some grains were found in the 1.2 A magnetic fraction. This procedure not only helps in gaining high quality heavy mineral concentrates, but also shortens the time of density separation using a heavy liquid. In this case, all obtained fractions were then separated using LST Fastfloat heteropolytungstate with a density of 2.9 g/cm³. After heavy liquid separation, the rutile grains were handpicked, mostly from the non-magnetic fraction and to a lesser extent from the 1.2 A fraction. Earlier investigations had shown that there is no correlation between the size of a detrital rutile grain and a magnetic fraction with its chemistry [46,47]; thus, the rutile grains were picked regardless of size or occurrence in magnetic fractions.

The grain surface morphology and initial mineral identification were performed using a ZEISS SIGMA VP electron microscope equipped with two energy-dispersive X-ray spectrometers Bruker XFlash 6/10 (SEM-EDS) in the National Multidisciplinary Laboratory of Functional Nanomaterials NanoFun, Faculty of Geology, University of Warsaw.
The selection of grains for further analysis was made using a stereoscopic microscope, taking into account the optical properties of rutile [48]. Subsequently, the rutile grains were mounted in epoxy resin and polished. Rutile compositions were determined by wavelength-dispersive spectrometry (EPMA) using a CAMECA SX Five field emission electron microprobe at the Joint-Institute Analytical Complex for Minerals and Synthetic Substances, University of Warsaw. The analyses were done under a 15 kV accelerating voltage, 75 nA beam current, and 5 µm beam spot size. The counting times 20 s at peak and 10 s for background for major elements were applied, while the intensity of Cr, Zr, and Nb analytical lines selected was measured for 40 s at the peak maximum, and the background intensity was recorded during half the time of the measurement at the peak position. The ‘PAP’ corrections after [49] were calculated. The standards supplied by Cameca and Structure Probe, Inc. were used during calibration of the microprobe. The standards, type of diffracting crystals, analytical lines used, and mean detection limits, expressed in ppm, were as follows: Mg (MgO, LTAP, Kα, 23), Al (Al2O3, TAP, Kα, 40), Si (SiO2, TAP, Kα, 12), Ti (TiO2, LLIF, Kα, 104), V (V, LLIF, Kα, 40), Cr (Cr2O3, LLIF, Kα, 37), Mn (rhodonite, LLIF, Kα, 20), Fe (hematite-SPI25, LLIF, Kα, 40), Zr (Zr, LPET, Lα, 20), and Nb (Nb, Lα, 20).

The EBSD (electron backscatter diffraction) technique [50,51] was used for identification of TiO2 polymorphs. Mounts with rutile crystals embedded in epoxy resin were polished using a 1.0 µm grain size diamond suspension and, furthermore, a 0.5 µm and 0.25 µm diamond suspension using a vibrating polisher (vibromet). Rutile grains, embedded in epoxy resin, were analyzed in Zeiss Auriga 60 FE-SEM (field emission scanning electron microscope) with a e− Flash Bruker EBSD detector. The rutile grains were not covered with a conductive carbon layer; however, for more effective electron removal, a selected area of the examined mount with rutile grains was framed with a highly conductive copper tape. The analyses were done in a high vacuum using a 10 kV acceleration voltage. The sample surface was tilted 70° relative to the beam and the detector tilt angle was close to 1.6°. The distance between the detector and the examined rutile surface was 17 mm. In most cases, rectangular areas within central parts of the analyzed grains were selected for EBSD analysis. Only an identification of TiO2 polymorphs was needed, therefore, a larger step size (pixel size of approximately 1.5-3.5 µm) than in standard EBSD crystallographic orientation studies was used. A phase map mode was used to identify common TiO2 polymorphs (rutile, brookite, or anatase) using the ESPRIT ver. 2.1 software, Bruker Nano GmbH, Berlin, Germany.

For TiO2 polymorph identification, the following crystallographic parameters, including space groups and fundamental unit cell parameters, were used: anatase (I41/amd; a = 3.7845 Å, c = 9.5143 Å), brookite (Pbca; a = 5.4558 Å, b = 9.1819 Å, c = 5.1429 Å), and rutile (P42/mnm; a = 4.5937 Å, c = 2.9587 Å).

4. Rutile Mineral Chemistry, Occurrence, and Zr-in-Rutile (ZIR) Geothermometry

Rutile is the most widespread and most thermodynamically stable natural polymorph of TiO2. The other, less common, natural polymorphs of TiO2 are akaogiite, anatase, brookite, and riebeckite [52,53]. Rutile most often crystallizes under conditions of medium- and high-grade metamorphism, and much less frequently in low-pressure hydrothermal quartz veins and in igneous rocks (for additional references, see Meinhold et al. and references therein [54]) with crystallization temperatures ranging from approximately 430 °C to over 1100 °C [31]. In general, rutile crystallizes commonly in metamorphic rocks formed under high-pressure conditions, i.e., above 1.1 GPa [54,55]. In rocks with a low Ca content, it is also stable at pressures lower than 0.7 GPa [56]. Low-pressure rutiles are also stable in low- and medium-grade metapelites, where the concentrations of Ca and Fe are very low and, conversely, the contents of Mg and Al are very high [54,55,57,58]. Low-pressure rutile may occur as characteristic needle-like crystals in quartz veins [59,60]. Taking into account the documented stability of rutile and the potential differentiation of the chemical compositions of the protolith rocks, detrital rutile occurring in sedimentary rocks most often crystallized in the temperature and pressure range corresponding to...
Rutile preserves information on the lithology of the source area, because of its very high resistance to weathering and diagenesis, whereas other minerals such as garnet, amphibole, and apatite may be completely dissolved occasionally. The use of rutile in provenance analysis is limited by the low concentrations of trace elements, which require the use of laser ablation inductively coupled plasma mass spectrometry (LA ICP-MS) or electron probe microanalysis (EPMA) in the trace analysis mode [31,33,47,54]. The use of EPMA in trace mode is facilitated by the high thermal stability of rutile, thanks to which it is possible to increase the detection limit of the determined trace elements by using the higher current of the electron beam and the extending count time at peak and background positions. Approximate concentrations of elements from the HFSE group (high field strength elements), represented by Nb, Ta, Zr, Hf, and W, as well as some transition metals group, such as Cr, V, and Fe, can reach concentrations up to 3000 ppm [30,33,54,63,64]. Zr concentrations in rutile from metamorphic rocks vary in the wide range from several ppm up to 11,000 ppm [29,34,65]. Anomalously high concentrations of transition metals are described only from rutile occurring in some ore deposits, e.g., [66], in which the contents of V$_2$O$_5$, Cr$_2$O$_3$, FeO, Nb$_2$O$_5$, and WO$_3$ can reach 3.95 wt.%, 3.28 wt.%, 3.27 wt.%, 0.23 wt.%, and 17.73 wt.%, respectively. The common high concentrations of HFSE and other transitional elements (particularly V and Cr) in rutile allow us to estimate rutile crystallization temperatures and discriminate between different source lithologies within source areas (for review, see [54]); both of these methods are commonly used in sediment provenance studies.

Zr-in-rutile geothermometry (ZIR) is a relatively new method and is used mainly for determining crystallization temperatures of rutile in various metamorphic and igneous rocks [67]. The key requirement for using the ZIR method is the determination of the Zr concentration in the rutile structure. Improving EPMA techniques in a trace mode allows us to determine the concentration of Zr with a detection limit below 40 ppm. The ZIR method can be applied only when the SiO$_2$–ZrO$_2$–TiO$_2$ system is in equilibrium with quartz, zircon, and rutile [31,32,65]. So far, in the literature, there are several calibrations of Zr in rutile geothermometers (Table 2). The first ZIR geothermometer (no 1 in Table 2) was proposed by [31] based on the Zr content of natural rutile crystals. This method assumes that, if rutile crystallizes in the presence of zircon and quartz, then Zr is buffered in the rutile structure. Rutile is able to preserve the original Zr concentration, indicating the peak of metamorphism even under retrogressive conditions [33], thus allowing for the determination of the relict temperatures of rutile formation from previous crystallization stages [68].

An alternative formula (no 2 in Table 2) was presented by [32], who calibrated their rutile geothermometer on experimental studies, supplementing it with only a few analyses of natural rutiles. The results from both geothermometers coincide only at about 540 °C. The discrepancy between the two geothermometers is clearly visible at lower and higher temperatures. This difference may come from the systematic analysis of rutiles only included inside garnets by [31], while rutiles found in the rock matrix were disregarded. The above mentioned geothermometers (no 1 and 2 in Table 2) are based only on the Zr concentration in the rutile structure. Geothermometers (no 3–7 in Table 2) presented by [65,69] additionally take into account the pressure values in the system during rutile crystallization. However, when analyzing rutile grains from sedimentary rocks, we have no certainty about pressure conditions occurring during their crystallization. In earlier studies, authors [47] suggested using a default 10 kbar and the α-quartz stability field geothermometer of [65] when information about pressure is not available. A comparison of calculated temperatures from the geothermometers of [32] (no 2 in Table 2) and [65] (no 3 in Table 2), with suggested default values, shows differences not greater than 5%. The results of both ZIR geothermometers are nearly the same; therefore, we chose not to make further
assumptions regarding pressure conditions and use the geothermometer calibrated by [32] (no 2 in Table 2). In the studies of detrital rutile performed by [28,29], it is claimed that more realistic results were obtained with the use of the [32] rather than the [31] geothermometer. Therefore, in this paper, as suggested by [28,54], only the thermometer proposed by [32] (no. 2 in Table 2) will be used.

Table 2. Compilation of the existing Zr-in-rutile (ZIR) geothermometers calibrated by various authors. The gas constant is \( R = 8.3145 \, \text{J} \cdot \text{mol}^{-1} \cdot \text{K}^{-1} \).

| No. | Formulae | Error of Temperature Estimation in [°C] | References for the Calibration |
|-----|----------|----------------------------------------|--------------------------------|
| 1   | \( T_{1}^{[\text{C}]} = 127.8 \times \ln\left(\frac{Zr_{[\text{ppm}]}^{K}}{4470}\right) - 10 \) | ±50 °C | [31] |
| 2   | \( T_{1}^{[\text{C}]} = \frac{7.36 - \log_{10}(Zr_{[\text{ppm}]}^{K})}{83.9 + 0.41 \times P_{[\text{bar}]}} + 273 \) | ±20 °C | [32] |
| 3   | \( T_{1}^{[\text{C}]} = \frac{0.1428 - R \times \ln(Zr_{[\text{ppm}]}^{K})}{1360 + 0.378 \times P_{[\text{bar}]}} - 273 \) | ±30 °C | [65] * |
| 4   | \( T_{1}^{[\text{C}]} = \frac{130.66 - R \times \ln(Zr_{[\text{ppm}]}^{K})}{57130 + 0.378 \times P_{[\text{bar}]}} - 273.1 \) | ±15 °C | [69] * |
| 5   | \( T_{1}^{[\text{C}]} = \frac{0.1433 - R \times \ln(Zr_{[\text{ppm}]}^{K})}{88.1 + 0.206 \times P_{[\text{bar}]}} - 273 \) | ±30 °C | [65] * |
| 6   | \( T_{1}^{[\text{C}]} = \frac{0.1412 - R \times \ln(Zr_{[\text{ppm}]}^{K})}{23910 + 0.249 \times P_{[\text{bar}]}} - 273.15 \) | ±30 °C | [65] * |
| 7   | \( T_{1}^{[\text{C}]} = \frac{139.63 - R \times \ln(Zr_{[\text{ppm}]}^{K})}{13000 + 0.249 \times P_{[\text{bar}]}} - 273.15 \) | ±15 °C | [69] * |

Note that the pressure units used in Formulas (4) and (7) are expressed in bars, while in Formulas (3), (5), and (6), they are in kilobars. *—equations for different quartz stability field.

The concentrations of other trace elements in the rutile structure can also be used to discriminate the source lithology. It was previously suggested [30,70] that differences in the Cr/Nb ratio may point to the metapelitic or metamafic origin. As mentioned earlier, the presence of quartz in the system is required for the correct use of the ZIR thermometer. In the case of mafic rocks, the presence of quartz may be the result of metamorphic reactions, even in undersaturated mafic protoliths [71–73]. Therefore, rutile derived from different lithologies can be successfully applied for calculating crystallization temperatures using a ZIR thermometer. Equation \( X = 5 \times (\text{Nb}_{[\text{ppm}]} - 500) - \text{Cr}_{[\text{ppm}]} \) proposed by [47] is considered as more accurate in discriminating source lithology using Cr/Nb ratios. However, the Nb-Cr discrimination method does not always give correct results [62,74]. The inaccuracies of the Nb-Cr discrimination may come from re-equilibration during retrograde metamorphism [74] or from usage of earlier, less reliable, and accurate discrimination equations [29,33,70]. In the case of crystalline rocks, the origin of rutile is generally known and a critical approach to Nb-Cr discrimination methods is possible. However, detrital rutile’s source rock crystallization conditions are usually unknown and, therefore, this discrimination technique must be applied with care and skepticism. Nevertheless, the Nb-Cr discrimination diagrams should be treated as an additional useful tool in studies of the origin of the clastic material [33,47,54].

5. Results
5.1. Mineralogy and Petrography of the Albian Sands

The Albian sediments are similar in all the examined sites throughout the Miechów Segment. They are composed of medium-grained quartz sands and sandstones devoid of calcium carbonates and phosphates, with variable admixtures of glauconite and muscovite. In Glanów-Stroniczki, the sample contains much higher contents of muscovite than in other outcrops; additionally, the largest crystals reach up to several mm in size. In most outcrops, apart from sands, there are also discontinuous, horizontally formed silification beds or irregular bodies of spongiolites. In the outcrop on the Chelmo Mount, the Albian sediments are lithified, medium-grained, and/or coarse-grained ferrigenous quartz sandstones. Quartz constitutes almost 90% of all detrital grains. The dominant quartz grains are subrounded and cemented by chalcedony and kaolinite. Potassium feldspar,
glaucnite, and heavy minerals are relatively rare. In sandstones from the Chelmo Mount, only a few, small, muscovite flakes were identified. The Albian sands at the Kopiec outcrop are largely similar to those found on the western side of the present-day HCM. These are clastic sediments dominated by medium- and coarse-grained quartz sands. In the sands of Kopiec, however, there are significant amounts of small pieces of fine-grained phosphorite rocks and carbonate cements (mostly calcite) that are not found in the Miechów Segment. The Albian sediments from the Łuków IG-1 and Piaski IG-2 boreholes are similar to each other. They constitute poorly cemented, fine-grained sandstones or loose quartz sands with a significant amount of glauconite and fine muscovite flakes. As in the sample from Kopiec, small grains/clasts of phosphorite are also present. The Albian sandstones from the Tomaszów Lubelski IG-1 borehole are poorly lithified, composed of medium-grained quartz with macroscopically visible glauconite. However, the concentrations of muscovite are much lower than in samples from the north (Łuków IG-1 and Piaski IG-2). The sample taken from the Potok IG-1 borehole is similar to that from Tomaszów Lubelski IG-1 and Kopiec; it is characterized by a significantly higher content of calcium carbonate and only a few broken muscovite grains were identified.

Although petrographic and mineralogical descriptions of the main mineral components of the Albian sediments from southern Poland are somewhat variable, the heavy mineral assemblage is exceptionally homogeneous and does not exceed 1% of the total grains of entire samples. In all of the examined samples, there is a dominance of minerals that are highly resistant to physical and chemical weathering. Thus, tourmaline, rutile, zircon, staurolite, and to lesser extent monazite-(Ce) dominate in heavy mineral assemblages. Garnet, spinel, kyanite, and sillimanite can also be found, but in much lower amounts. Significant and variable concentrations of Fe-Ti oxides, usually very strongly affected by leucoxenization, are also present in all of the analyzed samples.

The most frequent rutile grains are dark yellowish to reddish brown. The crystal size is variable and, on average, ranges from approximately 20 µm to 250 µm (Figure 3). Grains larger than 300 µm are rare and commonly rounded. Under the SEM and optical microscope, most rutile grains are euhedral, only partially angular, subangular, and subrounded in shape (Figure 3). Various degrees of abrasion can be distinguished on the grain surfaces. The edges and joints of the walls are often abraded to a higher degree than the wall surfaces. Rounded rutiles are subordinate (Figure 3a,g) and represent up to approximately 30% of all the rutile grains. Detrital rutile from the same outcrop or borehole shows all possible stages of abrasion and weathering, e.g., rutile grains from the Bolmin at the Miechów Segment (Figure 3b–d). Numerous grains show the pristine euhedral habit (Figure 3b); moreover, numerous grains have fresh cracks without weathering (Figure 3c), although highly altered grains are also common (Figure 3d). Rutile grains from the Łuków IG-1 borehole, the Puławy Segment, are also variable (Figure 3e,f). Some needle-like rutile crystals do not show any signal of alteration apart from initial physical weathering (Figure 3e), whereas other grains (Figure 3f) are altered (most likely by chemical etching) and then subrounded during physical weathering. The highest degrees of physical and chemical weathering are visible in rutile grains from the Tomaszów Lubelski IG-1 borehole, Puławy Segment. Numerous grains from this locality are highly abraded and chemically etched (Figure 3g,h), but as in the other localities, euhedral, pristine rutile grains can also be found.
Figure 3. SEM images of detrital rutile grains showing different stages of abrasion and weathering. (a) Rutile crystals from Glanów-Stroniczki (GLA) with visible chemical weathering marks; (b–d) grains from Bolmin (BOL) showing different stages of weathering from pristine crystals without traces of weathering (b); grain with fresh cracks without weathering (c); to highly altered crystal with visible etchings (d); (e,f) rutiles from borehole Łuków IG-1 (sample LK), needle-like rutile crystal (e) does not show any sign of alteration apart from physical weathering, (f) rutile is altered and rounded by weathering (chemical and physical); (g,h) altered and chemically weathered rutile from borehole Tomaszów Lubelski IG-1 (sample TL).

5.2. Results of Electron Backscattered Diffraction Analysis (EBSD)

To be able to correctly use the ZIR geothermometer, it is crucial to be certain that the mineral grains being analyzed are rutile and not the other common TiO$_2$ polymorphs such as anatase or brookite. When the ZIR geothermometer is applied to a TiO$_2$ polymorph other than rutile, the results may be highly inaccurate [47,75]. Because all the most common TiO$_2$ polymorphs, i.e., rutile, anatase, and brookite, have similar magnetic susceptibilities [45] and densities, it is difficult to separate them using only magnetic or density separation methods. The SEM-EDS method, as well as the electron probe microanalysis, are also not sufficient for accurate polymorph identification [75]. Other polymorphs, e.g., riesite and akaogiite [52,53], representing the high pressure varieties of TiO$_2$ polymorphs, are found in impact craters; however, they are extremely unlikely to be found in high abundances in clastic sediments. The correct polymorph identification of the rutile grains selected for chemical studies can be performed using Raman spectroscopy [76,77] or EBSD [51].
The electron backscattered diffraction analyses were done to check the precision of rutile separation from the Albian sands and to test for rutile identification on the basis of trace element concentrations as proposed by [75]. The results of EBSD phase mappings (Figure 4) do not show any polymorph other than rutile in the handpicked grain set, even in cases of uncertain grains. All analyzed TiO$_2$ minerals are rutile, and no other TiO$_2$ polymorphs such as anatase or brookite were recorded. The rutile grains do not contain any visible exsolution, twins, or crystallographically misoriented domains.

**Figure 4.** Results of electron backscattered diffraction analysis (EBSD) study of selected rutile grains. (a,b) Experimental EBSD Kikuchi patterns of rutile grains GLA-23 (a) and GLA-24 (b); (c,d) simulated Kikuchi pattern of the same rutile grains GLA-23 (c) and GLA-24 (d); (e) BSE (backscattered electron) image of rutile grains GLA 23 (top) and GLA-24 (bottom); (f) FSE (forescattered secondary electron) image of rutile grains GLA-23 and GLA-24 superimposed on EBSD mapping: red—rutile, single blue pixels*—anatase, single green pixels*—brookite; (g) BSE image of six more rutile grains MO-9—MO-14; (h) FSE (forescattered secondary electron) image of rutile grains MO-9—MO-14 superimposed on EBSD mapping: red—rutile, single blue pixels*—anatase, single green pixels*—brookite; *—single pixels of apparent anatase and brookite phase are most likely just a misidentification of the Bruker esprit software, Bruker Nano GmbH, Berlin, Germany used in EBSD study.

All TiO$_2$ grains with a very low V content (<50 ppm), which, according to the criteria of [75], might be classified as anatase, were also tested. The results of the EBSD analysis
showed that all the separated TiO$_2$ minerals with a low V content were rutile (see Figure 4, Table 3).

Table 3. Mineral chemistry of representative rutile from the Albian deposits of the SE Poland.

| Sample | GLA | GLA | KRZ | PRZ | PRZ | LEL | LEL | MO | MO | PO |
|--------|-----|-----|-----|-----|-----|-----|-----|----|----|----|
| No.    | 44219 | 44220 | 102/1 | 68/1 | 52/1 | 44203 | 34/1 | 72/1 | 77/1 | 44223 |
| Comment in wt.% | WG | WG | WG | WG | WG | WG | WC | WC | WC | WC |
| Ti     | 59.00 | 59.22 | 59.53 | 58.49 | 59.05 | 59.35 | 59.59 | 58.77 | 58.71 | 59.10 |
| Σtrace elements | 1.09 | 1.17 | 0.95 | 2.07 | 1.26 | 0.40 | 0.38 | 0.78 | 1.04 | 1.04 |
| O      | 39.87 | 40.07 | 40.12 | 39.99 | 40.02 | 39.80 | 39.94 | 39.60 | 39.66 | 39.88 |
| Total  | 99.96 | 100.45 | 100.61 | 100.54 | 100.32 | 99.56 | 99.88 | 99.14 | 99.41 | 100.02 |

| Source lithology | P | P | P | P | M | M | M | P | P | P |
| Temperature [°C] | 938 | 856 | 673 | 912 | 589 | 503 | 634 | 829 | 913 | 831 |

| Sample | BOL | BOL | GCH | TL | TL | PK | PK | LK | LK | KOP |
|--------|-----|-----|-----|----|----|----|----|----|----|----|
| No.    | 50/1 | 49/1 | 69/1 | 42/1 | 44227 | 44205 | 44209 | 61/1 | 48/1 | 44211 |
| Comment in wt.% | CG | CG | CG | EG | EG | EG | EG | EG | EG | EG |
| Ti     | 59.10 | 59.65 | 59.42 | 59.61 | 59.07 | 60.10 | 59.33 | 59.43 | 59.30 | 59.33 |
| Σtrace elements | 1.06 | 0.78 | 0.42 | 0.39 | 0.39 | 0.35 | 0.25 | 0.61 | 0.56 | 0.64 | 1.16 |
| O      | 39.88 | 40.11 | 39.88 | 39.99 | 39.99 | 39.58 | 40.25 | 39.89 | 39.91 | 39.87 | 40.01 |
| Total  | 100.03 | 100.54 | 99.83 | 99.98 | 98.99 | 100.50 | 99.82 | 99.90 | 99.82 | 100.50 |

| Source lithology | P | P | P | M | M | M | M | P | P | P |
| Temperature [°C] | 938 | 856 | 673 | 912 | 589 | 503 | 634 | 829 | 913 | 831 |

WG—western group rutile, CG—central group rutile, EG—eastern group rutile; * the source lithology was calculated using the equation proposed by [47]; ** crystallization temperatures were calculated after [32]; P—metapelite, M—metamafite; b.d.l.—concentration below detection limit.

5.3. Mineral Chemistry of Examined Rutile (EPMA)

The chemical compositions of 604 rutile grains, selected from the twelve localities of the examined Albian sands of SE Poland (see Figure 2), were determined by the electron microprobe method (EPMA). A representative data set is presented in Table 3, while the full dataset is attached to the paper as Supplementary Material (see Table S1). The analyses were considered valid when the total sum deviated by no more than 1.5 wt.% from 100 wt.%. Rutile analyses with an Si content higher than 200 ppm were also excluded from further consideration, as suggested by [31,54]. This step is necessary to rule out any rutile grains with zircon (ZrSiO$_4$) micro-inclusions, which may artificially increase the content of measured Zr, and thus inflate the calculated rutile crystallization temperature. Exsolution lamellae of baddeleyite (ZrO$_2$) were not found in any of the analyzed rutile grains.

The average concentrations of trace elements (Nb, Si, Zr, Al, V, Cr, Mg, Mn, and Fe) in rutile were within the ranges typical of different types of metamorphic rocks [30,54,55,62]. The sum of the content of these elements determined in the rutiles from the Albian deposits...
of SE Poland varied from 0.16 wt.% to 9.77 wt.% (see Table S1). More than 60% of the rutile analyses show the sum of the trace element content below 1 wt.%, while only 1.5% of the analyses have a total content of trace elements exceeding 3 wt.%. The measured Zr concentrations of analyzed rutile grains extend from 0 to 7490 ppm (see Table S1) with no distinctive internal chemical zonation. The range of Zr contents in nearly all samples varies from 4000 to over 7000 ppm. Only in the sample from the Łuków IG-1 borehole, the range is limited to 1020 ppm. However, the median Zr content within an analyzed sample rarely exceeds 1000 ppm. Rutile grains with very low Zr or with Zr below detection limits were also identified in all the studied samples (Table S1).

The concentrations of Fe (on average 1500–2000 ppm) and Cr (average 620–680 ppm) in rutile are uniform and do not vary significantly, both within and between samples. The mean Nb concentrations in most samples are above 1500 ppm. Only in Tomaszów Lubelski IG-1, Piaski IG-2, and Łuków IG-1, the mean Nb concentrations range from 740 to 830 ppm. Vanadium contents in rutile grains also show visible differences with regard to sampling location (see Table S1). In localities from the Miechów Segment, i.e., the geographically western zone (Przychody, Glanów-Stroniczki, Lełów, Korzkiew, see Figure 2) and central zone (Bolmin, Mokrzesz, Chelmo Mount, see Figure 2), the mean V concentrations are mostly above 1000 ppm. The highest concentrations of V, reaching over 13,330 ppm, are reported from Przychody (on average, in this sample, 8010 ppm) and Glanów-Stroniczki (average 5105 ppm). Rutile grains east of the HCM, i.e., from the Szydłowiec and the Pulawy Segments, which is the central zone (Potok IG-1) and the eastern zone (Tomaszów Lubelski IG-1, Łuków IG-1, Piaski IG-2, Kopiec, see Figure 2), contain much less V in their structure. Their mean V concentrations do not exceed 1000 ppm, e.g., in Kopiec, Tomaszów Lubelski-IG1, and Piaski IG-2, they range from 650 to 750 ppm. The lowest concentrations of V are in rutiles from Łuków IG-1, with a median value of less than 400 ppm (Table 3).

5.4. Estimation of Rutile Crystallization Temperature Using ZIR Geothermometer

Despite a significant range of zircon concentrations in each sample, distinctive peaks of estimated crystallization temperatures are indicated based on the rutile geothermometer calibrated by [32]. Estimates of the rutile grains are presented in Table 3 and Supplementary Table S1. Taking into account the temperature distributions in each of the tested samples (see Figure 5), three groups of samples (western, central, and eastern) were distinguished in Figure 6, which additionally allows us to divide the area of SE Poland into three regions (western zone, eastern zone, and central zone, see Figure 2), characterized by a similar spectrum of rutile crystallization temperatures.

The western group of samples includes four outcrops: Korzkiew, Glanów-Stroniczki, Przychody, and Lełów (Figure 2). The histograms of temperature distribution are poly-modal (Figures 5a and 6a). The majority of results oscillate around 700–800 °C; however, there are also visible peaks in the range of 550–600 °C as well as close to 900 °C. The distribution is platykurtic (Figure 6a) with a negative excess −0.6. Rutile compositions from the western group presented in the Nb-Cr diagram [47] show a significant dominant abundance of rutiles from matapelitic rocks over rutiles from metamafic rocks (Figure 7a). The content of metapelitic rutile grains varies between 80 and 90% of the population in each of the analyzed locations in the western zone, averaging 87% of metapelitic rutile, while the metamafic rutiles represent only 13%.

The eastern group also includes four sample localities, three boreholes: Łuków IG-1, Piaski IG-2, and Tomaszów Lubelski IG-1, as well as the Kopiec (Annopol) outcrop. All listed localities are positioned east of the Holy Cross Mountains (Figure 2). The temperature histograms are monomodal in this group of samples (Figures 5c and 6c). The distribution is leptokurtic (Figure 6c), with a positive excess of 6.7. The distribution of the results is also much more concentrated than that of the normal distribution. The rutile compositional data from the eastern part of the study area presented on the Nb-Cr discrimination diagram show a nearly even distribution of rutiles from metamafic and metapelitic rocks (Figure 7c). The content of metapelitic and metamafic rutile is similar, at 53% and 47% respectively. The
highest ratio of metamafic rutiles, reaching 57%, can be found only in the sample from the Piaski IG-2 borehole near Lublin.

![Figure 5. Histograms of rutile crystallization temperatures for all samples in the three groups: western (a), central (b), and eastern (c). Details of sample abbreviation and localization in Table 1. Temperatures were calculated using the [32] formula. Blue colour—metapelitic rutile; orange colour—metamafic rutile.]

![Figure 6. Summary histograms of rutile crystallization temperatures for the western (a), central (b), and eastern (c) groups with an indication of metamorphic facies. Temperatures were calculated using the Watson et al. [32] geothermometer.]

The content of metapelitic rutile grains varies between 80 and 90% of the population in each of the analyzed locations in the western zone, averaging 87% of metapelitic rutile, while the metamafic rutiles represent only 13%

Figure 7. Cr-Nb plots showing contrasting rutile provenance and pie charts of relative abundance of metamafic and metapelitic rutiles in the distinguished western (a), central (b), and eastern (c) groups. Discrimination of metamafic and metapelitic rutile is based on [47]. Details of sample abbreviations and localization are provided in Table 1.

The central group includes the following localities: Mokrzesz, Chełmo Mount, Bolmin, and borehole Potok IG-1 (Figure 2). The histograms of temperature distribution are bimodal in this group (Figures 5b and 6b). The one modal value varies around 550–600 °C, while the second is around 850–950 °C. The distribution is platykurtic (Figure 6b) with a clearly negative excess of −1.5. The rutile data from the central group presented in the Nb-Cr diagram show the prevalent abundance of rutiles from metapelitic rocks over rutiles from metamafic rocks (Figure 7b). The contents of metapelitic rutile vary on average around 83% of all the rutile grains, while metamafic rutile represents only 17%. The distribution of the
calculated crystallization temperatures, in both metamafic and metapelitic rutiles, does not differ significantly in all three groups (Figures 5 and 6).

6. Discussion
6.1. Geochemistry and Temperatures

The simple chemical composition of rutile (TiO$_2$) restricts the possibility of significant Ti substitution by other elements in the mineral structure [54]. This feature excludes rutile as a good tool for determining the provenance of clastic sediments solely on the basis of the variability in major element composition. It was suggested by [30] that rutile grains with more than 1000 ppm of Fe tend to be metamorphic in origin. This parameter is not as widely used as, e.g., in the Nb/Cr discrimination of lithology, but nevertheless, it can be useful in determining source rock lithology. In all the samples, the majority of rutile grains have more than 1000 ppm of Fe (Table S1, Supplementary Materials). In the western and central zones, they constitute c. 66.5%, while in the eastern zone, those metamorphic rutiles make up 80.1% of all analyzed rutile grains. Therefore, it seems that metamorphic rocks were the main sources for detrital rutile grains in Albian sands of southern Poland.

Rutile, in the petrographic study of high-grade metamorphic rocks, plays an important role as a constant component in the paragenesis of metamorphic rocks [51,54,55,57,62,78]. Petrological studies of igneous and metamorphic rocks have documented an increased concentration of trace elements such as Zr, W, Cr, V, Hf, Nb, Ta, and others [54,65,74,79]. These data are used to identify the protoliths of metamorphic rocks and in geothermometry and geobarometry [33,78,80]. The data obtained from PT studies of rutile crystallization conditions in metamorphic and igneous rocks have also been successfully applied to detrital rutile [28,29,33,34,81].

Estimation of rutile crystallization temperatures using a ZIR geothermometer [31,32] gives a wide range of results ranging from <500 °C to over 1000 °C (Figures 5 and 6, and Table S1). The estimated crystallization temperatures of rutile in the Albian sand samples that were collected from outcrops (Table 1) in SE Poland point to metamorphic facies: greenschist–blueschist (<550 °C), amphibolite–lower temperature (LT) eclogite (550–700 °C), and granulite–high temperature (HT) eclogite (above 700 °C) (Figure 8).

![Figure 8](image-url)  
**Figure 8.** Pie chart of relative abundances of rutiles from different metamorphic facies in the western, central, and eastern groups.
Platykurtic distribution of calculated rutile temperatures from the western and central groups indicates rather variable sources of the detrital material. Rutile with temperatures peaking at 700–800 °C and above 900 °C was derived mainly from granulite–HT eclogite facies rocks (65% in the western group and 52% in the central group), while a much smaller amount of rutile grains with temperatures peaks at 550–600 °C were probably derived from amphibolite–eclogite facies rocks (32% in the western group and 35% in central group). Single grains (3% in the western group) and non-common grains (13% in the central group) were derived from greenschist–blueschist facies rocks (Figure 8). By contrast, monomodal distribution of temperatures in the eastern group points to one source rock and a common area of provenance of the rutile grains. Most results vary around 550–600 °C and are derived from amphibolite–LT eclogite facies rocks (66%).

Considering the distribution of rutile crystallization temperatures within the three groups, the eastern one is clearly characterized by rutile with crystallization temperatures peaking at 500–600 °C (Figure 6), while such rutile occurs in only small amounts in the western group. In turn, rutile with a crystallization temperature of 700–800 °C dominates in the western group, but appears sporadically in the eastern group. Such characteristics clearly indicate at least two different source areas providing material for the detrital rutile. This is confirmed by the proportion of rutile originating from metapelitic and metamafic rocks. In the western group, metapelitic rutile grains clearly dominate over metamafic rutile, while in the eastern group, the amounts of metapelitic and metamafic rutile are nearly equal (Figure 7). Additionally, the V contents in rutile also point to different sources for the western and eastern zones. On average, rutiles grains from the western zone contain c. 1910 ppm of V, which contrast with contents around 625 ppm in the eastern zone (Table S1). The detrital rutile of the western and eastern groups appears to come from various host rocks and probably from different source areas.

Interesting data come from the central rutile group (Figure 6). In this group, the amount of rutile with temperatures of crystallization in the range of 500–600 °C is clearly visible, which are those that dominate the eastern group and are of minor importance in the western group. So, here we have a clear influence of detrital rutile from the same source as rutile in the eastern group. On the other hand, rutile with the crystallization temperature range of 700–800 °C in the central group is present, but is not so dominant as in the western group. Moreover, in the central group, there is a distinct group of rutile grains with crystallization temperatures ranging from 800 °C to 950 °C, which are also present in smaller amounts in the western group, and very sporadically in the eastern group (only five grains). Thus, such data may indicate that part of the detrital material was supplied to the central zone from the same source as the western zone. The last interpretation is confirmed by a very similar proportion of metapelitic rutile to metamafic rutile in the central and western groups (Figure 7). It seems, therefore, that the group distinguished as central was influenced by various lithological sources and probably provenance. The detrital material deposited in the central zone was derived from the same source as in the western group, but also has some signs of the same source as that of the eastern group, i.e., the central zone constitutes a distinct mixed group. Additionally, this group has a clear large admixture from high-temperature granulite and HT eclogite facies rocks (above 850 °C) similar to the western group.

A multidimensional scaling (MDS) statistical analysis was used to compare the similarities between the samples. The MDS is a technique that compares large datasets of a variable, e.g., temperature, element concentration, detrital age, and so on [82]. The result of this method is a configuration of points in the two-dimensional Euclidean plane, creating some sort of a map in which samples with similar analyzed variable values tend to cluster close to each other, while samples with greater dissimilarities plot further apart [83–86]. It is worth noting that the MDS diagrams of calculated rutile crystallization temperatures and Nb concentrations visualizing the level of similarity between samples confirm quite well the division into three groups adopted by us based on the ZIR method (Figure 9). The four localities of the eastern zone (samples TL, PK, LK, and KOP) on the MDS diagrams are...
very close to each other, which indicates a high statistical similarity between those samples. The three samples from the central zone (MO, BOL, and GCH) are also located together on the temperature MDS diagram (Figure 9a), which confirms their statistical similarity. One sample, PO, is located between the TL, PK, LK, KOP and MO, BOL, GCH sample groups. Geographically, the PO sample is close to the eastern zone; however, temperature data allow us to include that sample in the central zone (Figure 5). The four samples from the western zone (GLA, PRZ, LEL, and KRZ) are close to each other on the temperature MDS diagram, reinforcing their inclusion in the same, western group. Similar conclusions can be derived from the MDS diagram of Nb concentrations (Figure 9b). As in the temperature MDS diagram, the eastern group is clearly distinct from the rest of the samples. However, the samples of the central group are not as clearly separated from the western group as shown in the temperature MDS diagram. Nonetheless, a faint division between these two groups can still be distinguished.

**Figure 9.** Nonmetric multidimensional scaling (MDS) maps of (a) calculated crystallization temperature of detrital rutile grains using the Watson et al. formula [32] detailed explanation in the text; (b) Nb (ppm) concentrations in rutile.

### 6.2. Source Areas and the Paleogeography

The next question is about the source area(s) for sands of the extra-Carpathian of southern Poland during the Late Albian. The answer to this question requires a combination of our results with the paleogeography studies of central Europe (Figure 1). During the Neo-Cimmerian phase, in the early Cretaceous, a significant regional uplift of some areas of southern and western Europe with numerous block tectonic movements took place [87]. In the Middle and Late Albian (110.8–100.5 Ma) during the Cretaceous eustatic transgression [87,88], the sea encroached onto a denuded area of central and southern Poland [35,41]. This part of Poland covered by the epicontinental sea (Figure 1) was the easternmost part of the Central European Basin System [87]. The transgression encroached onto the Polish Lowlands roughly from the NW direction using an earlier tectonic setting of the Polish Trough, and propagated in a south-southeasterly direction.

The area of central and southern Poland was covered with various types of Upper Jurassic carbonate deposits and siliciclastic rocks of the Lower Cretaceous [8,35,42]. The transition between the Albian and Jurassic carbonate sediments is erosional, and basal conglomerates can often be observed [35,89]. In some locations, the top of the Jurassic carbonates shows a distinct silification and karst surfaces that point to the regional terrestrial character of the pre-Albian relief [89,90], which is supported by recognition of backshore settings during the Late Albian in the western part of the Miechów Segment [91].
Jurassic dark flints found in the Albian sands in the Miechów segment additionally indicate erosion of the Jurassic carbonate sediments during the transgression [89]. During the Albian, in the area of the contemporary Miechów Segment in the western part of the study area, a large peninsula extending NW-SE was located, reaching its end near Radomsko and continuing south beyond the Carpathian thrusts [92,93]. This peninsula is devoid of Albian sediments and the Cretaceous sea transgression covered that area in the Cenomanian and/or Turonian. The Upper Jurassic limestone is covered in some places directly by Cenomanian calcareous sandstone (e.g., Zajęcza Góra near Skotniki) of the northeastern part of the Miechów Segment [94]. In the southern part of the Miechów Segment, e.g., in the Wielkanoc Quarry, the Turonian limestone directly overlies the bihermal Upper Jurassic limestone [42,95]. The area of the Sudečes was also not flooded by the sea in the Albian, but remained an elevated land area until the Cenomanian, when the transgression began to invade this area as well [94,96]. Further to the east of the present-day HCM, the Albian rocks are covered with carbonate sediments of the Upper Cretaceous [97]. The Albian deposits have been drilled in the western and northern parts of the Lublin region. Albian deposits are represented here by fine and medium-grained quartz sands and/or weakly lithified sandstone, changing up in the section into sands/sandstone with admixture of calcium carbonate and marls with numerous phosphate rocks [98]. The Albian deposits do not occur close to the eastern border of Poland and on the SW of Lublin towards Kraśnik [93,97]. Here, the Cenomanian or Turonian carbonate sediments of Upper Cretaceous directly overlie the older Pre-Cretaceous sediments [97]. The geological data mentioned above suggest the presence of a developed coastline with distinct sea bays and with the dynamic shifting the coastline during the transgression in the study area. The sea in Albian times encroached over an area covered by sedimentary rocks, mostly carbonate and to a lesser extent siliciclastic rocks. Part of the clastic material found in the Albian sediments could thus have come from the recycling of older sedimentary rocks, most likely Lower Cretaceous sediments. Those sedimentary rocks cannot have been the primary source of rutile. The sources of this mineral can be found mainly in metamorphic or igneous rocks. Land massifs of crystalline rocks did not occur in central and northern Poland in the early Cretaceous. The probable source areas of clastic material during the Albian are the closest lands of crystalline massifs such as the Bohemian Massif to the west/south-west, the Ukrainian Shield in the east, and the Baltic Shield to the north of the study area (Figure 1).

The latest geochemical studies on tourmaline and monazite from Albian sands of the Miechów Segment point to the Bohemian Massif as the most probable source area [25,99]. The Bohemian Massif comprises a large variety of igneous and metamorphic rocks [100,101]. Therefore, it is possible to indicate potential source rocks for detrital rutile crystallized under amphibolite–LT eclogite facies (e.g., mica schists, gneisses) conditions [100,102,103], as well as in high-grade granulite–HT eclogite facies rocks, namely granulites and gneisses [101,104,105]. During the Albian, the Bohemian Massif was neighboring to the western and central zones (Figure 2). Moreover, the study of heavy mineral provenance of Jurassic and Triassic sedimentary rocks in southern Poland also points to the Bohemian Massif as a main source of detrital material [106,107]. It is thus most certain that the detrital heavy minerals found in the western group and three samples from the central group (MO, GCH, and BOL) from the Miechów Segment have come from the Bohemian Massif.

The Bohemian Massif is the most probable source area for the western group of rutiles, but a more enigmatic transport direction of clastic material should not be disregarded. Southward from the study area, obscured by Neogene Carpathian thrusts, crystalline Silesian and Proto-Silesian Ridges were located. The ridges are composed mostly of granites, gneisses, schists, and quartzites [21,108]. The ridges are overlain by Carpathian thrusts, and thus information on their petrology and mineralogy comes from pebbles and exotics found in Carpathian sediments. Precise information on the P-T conditions of these rocks is not available; however, some data from garnets suggest high-grade metamorphic conditions, similar to those found in Moldanubicum and the western part of the Bohemian Massif [21,109].
As discussed earlier, the western and eastern groups had different sources of detrital rutile. The Bohemian Massif probably should not be considered as a primary and main source area for the eastern group. That group roughly corresponds to the Lublin area in eastern Poland (the eastern zone) and is situated approximately at the same distance between the Bohemian Massif and the Ukrainian Shield. The crystalline rocks in the Ukrainian Shield (which were most likely exposed during the Albian) consist mostly of Paleoproterozoic and Archean rocks such as granites, gneisses, TTG (Tonalite-Trondhjemite-Granodiorite), various mafic, and ultramafic rocks [102,110–112] that mostly underwent high-grade metamorphic conditions. Crystalline rocks found in the Podolia domain (southwestern part of Ukrainian Shield) are represented mostly by a high-grade granulite facies, granites, and gneisses [113]. High-grade metamorphic rocks (above 700 °C) are a highly unlikely source for the amphibolite–LT eclogite facies (550–700 °C) rutile occurring frequently in the eastern group in Albian sands. A wide range of various mafic and ultramafic rocks (such as pyroxenites, amphibolites, and anorthosites) in the Ukrainian Shield also could not have been the main source because of the scarcity (or absence) of rutile in their mineral assemblage [114]. The Neoproterozoic basalts and tuffs of the Volynian Flood Basalts province covering a vast area of eastern Poland, western Ukraine, southwestern Belarus, and northern Moldavia [115] are also an unlikely source of rutile. This is owing to a relative rarity and a small grain size (few micrometers, often less than 10 µm in diameter) of rutile found in effusive rocks [116]. The detrital rutile from the Albian sands in this work is much larger (20 µm to 250 µm, Figure 3). Thus, it is difficult to find the probable source area in an easterly direction (the Ukrainian Shield) for Albian rutile from the eastern group. It cannot be excluded that unknown source rocks have been completely eroded from the Ukrainian Shield.

The other possible source area for the detrital rutile of the eastern group may be the Baltic Shield located north of the study area. The Baltic Shield was located almost twice as far as the Ukrainian Shield and the Bohemian Massif during Albian times. The central and southern parts of the Baltic Shield are dominated by Palaeoproterozoic and Archean orogenic belts, while the Scandinavian Caledonides occur solely in the western part [110,117,118]. Calculated P-T paths for various rocks in the Scandinavian Caledonides show similar crystallization temperatures to the calculated rutile crystallization temperatures from the study area [119–121]. Analyses carried out on Albian detrital rutile from sandstone sediments of the Norwegian sea showed good convergence for the calculated crystallization temperatures (peaking at~550–600 °C) to our rutile from the eastern group, and in the proportion of metamafic to metapelitic rutile. This proportion in sandy sediments of the Albian MN3 sand types in the Norwegian sea is 60% to 40% [28], whereas in samples from eastern Poland studied here, it is c. fifty–fifty. It is possible that rutile has been derived from western Norway, from the Western Gneiss region and the Middle Allochton of the Caledonian Nappe Domain, where high-grade mafic gneisses, eclogites with pyroxenites, and peridotites are located [110,118,121–124] These regions sent material to the Norwegian Sea prior to the final opening of the northeast Atlantic [28,125]. It is also possible that this region, with its vast area of metamorphic rocks, was supplying detrital material to the epicontinental sea southward to the Mid-Polish Trough. The hypothetical longshore currents and grain shore drift distributed the clastic material (including rutile) from NW to SE.

Longshore currents can transport loose sediments along a coast parallel to the shoreline. The combination of longshore drift (beach drift) and longshore currents leads to longshore transport of detrital material [126]. It is possible that such transport took place along the NE shore of the elongated shallow Mid-Polish Trough. Despite quite long distances, about or greater than 1000 km, rutile is a very stable mineral in sedimentary environments and can be transported on long distances [27,127–129]. The transport of detrital material by longshore currents is common in recent seas where the combination of wind, wave action, morphology of the sea bottom, and sea-tidal action allows for the creation of such currents [130–132]. The Orange River carried diamonds from highlands
around Kimberly, South Africa, westward to the beaches of the South Atlantic Ocean. As the Atlantic gradually opened, the river deposited much of its diamond-bearing sediment on the Namibian coast and longshore currents carried them northward along the 600 km coast along the Namibian shore, and that transport distance is unequivocally documented with the diamond placer deposits [133]. We have not found information in the literature as to how far single diamond grains were transported near the Namibian shore towards the north, but from the mouth of the Orange River, sand material was transported alongshore for >700 km [128]. The longshore current transported the sediment material in the Mediterranean Sea from the Nile Delta, Egypt, to Haifa, Israel, a distance of over 450 km [134]. In the Baltic sea, longshore currents are generated along the Polish sand shore [126] and can transport sandy material along the entire southern Baltic shore [135,136]. Therefore, material transport by longshore currents over a distance of 1000 km is possible and plausible.

The Nb/Cr ratios and temperatures of crystallization of rutile from the central group (samples MO, GCH, BO, and PO) show their similarity to the western group; thus, the Bohemian Massif seems to be the main source for both groups. This is confirmed by studies of tourmaline and monazite from the MO, GCH, and BO samples collected from the Miechów Synclinorium [25,99]. On the other hand, the increased amount of rutile with crystallization temperatures in the range of 500–600 °C documented in the central group (Figure 6) shows their similarity to the eastern group of rutile grains. Hence, this component documents the significant proportion of detrital material transported probably from the Baltic Shield as in the case of the eastern group. During the Albian, the central zone (see Figure 2) can be interpreted as a mixing zone to which detrital material containing rutile representing the western and eastern groups was delivered.

The absence of high-temperature rutile in the Lublin area (the eastern zone) shows that detrital material derived from the Bohemian Massif does not exceed the contemporary HCDML land area (Figure 1). The distance between the western and eastern zones is not large enough (approximately 250 km) to easily explain the lack of “Bohemian rutiles” in the eastern zone. As mentioned above, rutile can be transported over much longer distances [27,127,129]. It seems, therefore, that there must have been orographic or bathymetric barrier(s?) in the study area between the western and eastern zones; however, the distribution of a sandy Albian facies near the contemporary HCDML does not unequivocally confirm the existence of a land area there [6,9,12,41,92,93]. Thus, the large peninsula extending NW–SE in the middle part of the present-day Miechów Segment postulated by Hakenberg [35] could have played the role of such an orographic barrier controlling the transport of detrital material from the crystalline Bohemian Massif to the western and partly central zones of the study area (Figure 2). Longshore currents along the southern shore of the Albian epicontinental sea could also have played an important role in the distribution of detrital material including the heavy minerals. The detrital rutile grains were transported to the area referred to by us as the central zone from the Bohemian Massif (from the west) and possibly from the southern part of the Baltic Shield from the north. During the Albian, between the Lublin area and the Ukrainian Shield, a bathymetric barrier [137,138] could have existed; there was no possibility for any longshore action and eastward and/or westward transport of detrital material into the study area.

7. Conclusions

The amount of accessory heavy minerals is less than 1% in predominantly quartz sand samples of the Albian sediments from SE Poland. The chemically and physically resistant rutile reach up to 20% of the heavy mineral assemblage in the Albian quartz sands of southern extra-Carpathian Poland.

Rutile grains, selected for EPMA study, are represented solely by the rutile polymorph that allows the usage of the Zr-in-rutile (ZIR) method to determine the rutile crystallization temperatures. Three groups of samples with estimated crystallization temperatures were distinguished in the studied material: western, central, and eastern.
The western group, which includes outcrops situated in the southwestern part of the Miechów Segment (western zone), indicates rather variable sources of detrital material, but mainly derived from metapelitic rocks. The detrital rutile grains were mainly derived from granulite–HT eclogite facies rocks (above 700 °C) with much smaller amounts of rutile grains derived from amphibolite–LT eclogite facies rocks (550–700 °C). The most likely source area for that group of rutile seems to be the Bohemian Massif. The Bohemian Massif has a vast spectrum of metamorphic rocks with various grade of metamorphism, thus it is a good source for the diverse crystallization temperatures of the detrital rutile in the western group.

The eastern group includes sampled localities situated northeast from the Holy Cross Mountains in Lublin area (eastern zone). Detrital rutiles were mainly derived from amphibolite–LT eclogite facies of metapelitic and metamafic rocks in similar quantities. The detrital rutiles were derived from a little petrographically diversified source area. The most likely provenance for that group of rutile seems to be the southern part of the crystalline Baltic Shield, possibly from the Western Gneiss region and the Middle Allochton of the Caledonian Nappe Domain. The hypothetical longshore current was flowing in a NW to SE direction along the NE nearshore zone of the Mid-Polish Trough and transported detrital material, including rutile.

The central group includes sampled localities situated in the northern part of the Miechów Segment and one sampled place (Potok IG-1 borehole) situated northeast from the Holy Cross Mountains (central zone). Detrital rutiles from this group were mainly derived from granulite–HT eclogite facies rocks and amphibolite–LT eclogite facies rocks and chiefly from metapelitic rocks. The source of rutiles was diverse, as in the western zone. Thus, the most likely source area for that group of rutile seems also to be the Bohemian Massif and, probably, to a lesser extent, the Baltic Shield.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/min11060553/s1, Table S1: Chemical composition of rutile from the Albian sands in Southern Poland.

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