Micro-Raman Spectroscopy of Selected Macerals of the Huminite Group: An Example from the Szczerców Lignite Deposit (Central Poland)

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Abstract: Lignite (ulminite reflectance Rr = 0.27%) from the Szczerców deposit (Central Poland) is dominated by huminite group macerals, containing a high proportion of attrinite and densinite. Densinite and ulminite are more abundant in small aromatic units than attrinite, which may result from their stronger gelification. The differences in Raman spectral characteristics between attrinite and ulminite are more pronounced than between attrinite and densinite. Fusinite, in comparison with the huminite group macerals, is composed of larger, more varied aromatic systems. The D4 (1190–1200 cm\(^{-1}\)) and D5 bands (1280–1290 cm\(^{-1}\)), most likely, correspond to different chemical structures, and their origin should be further investigated.

Keywords: lignite; Raman spectroscopy; huminite; attrinite; ulminite; densinite

1. Introduction

Lignite is an important energy carrier in many countries. Its chemical and technological features, which depend to a large extent on the petrographic composition, have been intensively studied for many years [1–4]. However, the microstructure of lignite macerals is still poorly known. So far, Raman spectroscopy studies aimed at understanding brown coal’s microstructure have been conducted almost exclusively on bulk samples [5–11].

The purpose of this work was to reveal, for the first time, and compare the microstructural features of attrinite, densinite, ulminite, and, additionally, fusinite, based on the example of the Szczerców deposit. It is one of two deposits currently exploited by the Polska Grupa Energetczna (PGE) open-pit mine Belchatów—the largest lignite mine in Europe.

2. Geological Setting

The Szczerców lignite deposit is located in the Szczecin–Łódź–Miechów synclinorium, which is located in the western part of the Kleszczów Graben [12] (Figure 1). The Szczerców deposit is approximately 7 km long and 1.5 to 3 km wide [13]. From the east, it is adjacent to the Belchatów lignite deposit. They are separated by the Dębina salt dome [13]. The Szczerców lignite overlies Upper Jurassic, Upper Cretaceous, and, locally, Paleogene sediments. The coal-bearing sediment is divided into [13]: a sub-coal complex, a coal complex, a clay and coal complex, and a clay–sand complex (Figure 1). The thickness of the coal complex reaches up to 100 m in the central part of the deposit; the average thickness is approximately 30–40 m [14,15].

The main seam consists of detritic and xylodetritic coal. The average content of xylites is approximately 20%. Occasionally, paralonstein (clay kaolinite) interbeddings are observed [14]. The lower part of the seam is of Lower Miocene age. Stratigraphically, it is
part of the lowest part of the Ścinawa Formation [16]. The hanging wall part of the seam is of Middle Miocene age [17]. Quaternary deposits are glacial, interglacial, and sandy sediments with lenses of gravels and glacial clays [13,18].

Figure 1. Location of Szczerców deposits (a), tectonic map of the deposit (b), and geological cross-sections (c) according to Czarnecki et al. [13] and Kasiński et al. [19]; modified.

3. Methods

The sample for microscopic and Raman spectroscopic investigation was collected as a representative channel sample from the seam in the western part of the Szczerców deposit. Two polished sections—one for each type of examination—were prepared according to the ISO 7404-2:2009 standard [20]. Petrographic analysis was conducted under incident white light and blue light excitation (i.e., fluorescence mode) using a Zeiss Axioplan polarizing microscope.

The random reflectance of ulminite B and other macerals was determined using monochromatic light with a wavelength of 546 nm and immersion oil ($n = 1.518$), and a Zeiss MPM-400 reflectometer according to the ISO 7404-5:2009 standard [21]. The petrographic composition of coal was determined according to ICCP guidelines [22–24]. The analysis was performed in accordance with the ISO 7404-3:2009 standard [25].

In addition, proximate and ultimate analyses were performed in an accredited laboratory in accordance with applicable standards for total moisture [26], ash content [27], volatile matter content [28], sulfur content [29], gross calorific value [30], and carbon and hydrogen content [31].

Raman spectroscopy of attrinite, densinite, ulminite, and fusinite macerals was performed in 27 to 30 randomly chosen points for each maceral. It should be noted that Raman spectra measurements were made on pure macerals. The selected measurement points are shown in the photographs in Appendix A (Figures A1–A4). The spectra were recorded with a Thermo Scientific DXR Raman (Faculty of Geology, Geophysics and Environment
Protection, AGH University of Science and Technology, Kraków, Poland) microscope with a 900 grooves/mm grating and a Charge Coupled Device (CCD) detector. The Olympus 10× (NA 0.25) objectives (spot sizes 2.1 µm and 1.1 µm, respectively) were used. Excitation was activated with a 532 nm diode laser with a maximum power of 10 mW. Measurements were performed in a spectral range of 400–3500 cm⁻¹, at a spectral resolution of 1 cm⁻¹, and an area of 1 µm². The laser power was set at 1–2 mW. The spectrometer was calibrated using a polystyrene standard. The accumulated measurement time was 30 s for each spectrum. Peak-fitting was conducted in the range of 1000–1800 cm⁻¹ by GRAMS32, based on the earlier Raman studies of low-rank coal (brown coals) [6–11]). The second derivative of the spectra was also considered to find the initial positions of the Raman bands. Lorentzian curves were applied. The goodness of fit was checked by the χ² test. The I_D1/I_G ratio was determined from the D1 and G band intensities. Furthermore, the A_D3 + A_D6/A_ALL, A_D4/A_ALL, and A_D5/A_ALL ratios were calculated from the band areas, where “A_ALL” denotes the sum of all band areas in a given spectrum. Raman band separation (RBS)—i.e., the distance between the G and D1 peak—was also determined.

The nomenclature of the bands was taken from Henry et al. [32]. The additional band D7 (~1095 cm⁻¹), which was earlier found by Li et al. [6] and labeled as S_R, was introduced.

The results of attrinite examinations were compared with those of densinite, ulminite, and fusinite using Student’s t-test with a significance level of α = 0.05. This was preceded by the Shapiro–Wilk test to check the normality of distribution, and the Fisher–Snedecor test to assess the equality of variances.

4. Results and Discussion

4.1. Petrographic Composition

Based on the macroscopic analysis (Table 1), the tested coal is xylodetrictic coal with a xylite content of 22%.

Table 1. Petrographic composition of lignite from the Szczerców deposit.

| Component       | Content (%) |
|-----------------|-------------|
| **Huminite**    |             |
| Textinite       | 11.7        |
| Ulminite        | 12.1        |
| Attrinite       | 29.9        |
| Densinite       | 23.9        |
| Corpohuminite   | 1.6         |
| Gelinite        | 1.5         |
| **Macerals**    |             |
| Sporinite       | 0.6         |
| Cutinite        | 0.1         |
| Resinite        | 1.1         |
| Suberinite      | 0.2         |
| Alginite        | 0.1         |
| Liptodetrinite  | 3.6         |
| **Liptinite**   |             |
| Fusinite        | 1           |
| Semifusinite    | 0.2         |
| Funginite       | 0.1         |
| Micrinite       | 0.0         |
| Inertodetrinite | 1.8         |
| **Inertinite**  |             |
| Pyrite          | 0.8         |
| Carbonates      | 0.5         |
| Quartz + Clays  | 9.2         |

The coal from the Szczerców deposit is dominated by macerals of the huminite group (80.7% vol.). The huminite macerals are dominated by attrinite (29.9% vol.) and densinite (23.9% vol.). The sum of these macerals indicates the dominant share of the detritic component in coal. Petrographic components of woody remnants, namely textinite
(11.7% vol.) and ulminite (12.1% vol.), were observed. The huminite components of the tested coal also include gelinite and corpohuminite (1.5 and 1.6% vol., respectively) (Table 1).

The content of liptinite macerals is low (on average 5.7% vol.). The most commonly observed component of this group is liptodetrinite (3.6% vol.). Terpene resinite (on average 1.1% vol.) is mainly an impregnation of sapropelic xylites occurring in a subordinate amount. The content of sporinite is 0.6% vol. The content of cutinite (0.1% vol.), alginite (0.1% vol.), and suberinite (0.2% vol.) is very low.

The content of inertinite macerals is 3.1% vol. The most abundant component in this maceral group is inertodetrinite (1.8% vol.), randomly dispersed in attrinite and densinite. Less-abundant elements of this group include fusinite (1.0% vol.) semifusinite (0.2% vol.), and funginite (0.1% vol.).

The mineral matter content of the examined coal is 10.5% vol. (dry basis. The most commonly observed minerals are clay and quartz, which are dispersed among attrinite and densinite. The content of these minerals is 9.2% by volume. Pyrite, which accounts for 0.8% by volume, is less abundant, and is most commonly found in the form of framboidal pyrite. The presence of carbonates, which are associated with lake chalk, is typical for coal from the Szczerców deposit [14]. The mineral matter is dispersed in the macerals and locally impregnates intracellular spaces. Dispersed clay minerals are most commonly observed in attrinite.

4.2. Random Reflectance and Proximate and Ultimate Parameters of Lignite from the Szczerców Deposit

The random reflectance was measured on five macerals according to the ISO 7404-5 standard [21]. The measurements on huminite macerals were made on textinite, ulminite, attrinite, and densinite. In the case of macerals of the inertinite group, the random reflectance was measured on fusinite (Table 2). The obtained results correspond well with the results obtained by Sýkorová et al. (2005). The huminite macerals textinite (0.2%) and densinite (0.31%) have the lowest and highest random reflectance, respectively. Ulminite B’s random reflectance is 0.27%.

Table 2. Random reflectance of selected macerals in lignite from the Szczerców deposit.

| Maceral     | Random Reflectance (%) | Standard Deviation (%) |
|-------------|------------------------|------------------------|
| Textinite   | 0.20                   | 0.034                  |
| Ulminite    | 0.27                   | 0.056                  |
| Attrinite   | 0.26                   | 0.039                  |
| Densinite   | 0.31                   | 0.060                  |
| Fusinite    | 0.95                   | 0.067                  |

According to the ISO 11760:2005 [33] standard and in-seam coal classification [34], the investigated lignite was low-rank coal (lignite C)—as the average reflectance of ulminite B is less than 0.40% (R<sub>r</sub> = 0.27%), the total moisture of the coal is >35% (51.3%), and the ash-free moisture is less than 75% (43.8%). The results obtained are consistent with other chemical measures of the rank of coal (C<sub>daf</sub> = 67.39%) and the gross calorific value (as received) (GCV<sub>maf</sub> = 13.9 KJ/kg) (Table 3), which classifies the examined coal as ortholignite.
Table 3. Proximate and ultimate analysis of lignite from the Szczerców deposit.

| Proximate Analysis                                      |   |
|---------------------------------------------------------|---|
| Total moisture, as received basis, M<sub>tr</sub> (%)   | 51.3 |
| Total moisture, moisture, ash free basis of M<sub>maf</sub> (%) | 43.8 |
| Ash, dry basis A<sub>db</sub> (%)                       | 18.6 |
| Total sulphur, dry basis, S<sub>db</sub> (%)           | 3.89 |
| Gross calorific value, moisture, ash free basis GCV<sub>maf</sub> KJ/kg | 13.9 |
| Gross calorific value, dry, ash free basis GCV<sub>daf</sub> KJ/kg | 24.70 |
| Volatile matter, dry, ash free basis V<sub>daf</sub> (%) | 49.37 |

Ultimate analysis of lignite

|                                |   |
|--------------------------------|---|
| Carbon content, dry, ash free basis C<sub>daf</sub> (%) | 67.39 |
| Hydrogen content, dry, ash free basis H<sub>daf</sub> (%) | 5.18 |
| Nitrogen content, dry, ash free basis N<sub>daf</sub> (%) | 0.89 |
| Oxygen content, dry, ash free basis O<sub>daf</sub> (%) | 21.47 |

db—dry basis; maf—moisture, ash free basis; as—as received basis; daf—dry, ash free basis.

4.3. Micro-Raman Spectroscopy

4.3.1. General Characteristics of the Raman Spectra

Representative Raman spectra of attrinite, densinite, ulminite, and fusinite from the studied coal are given in Figure 2. Eight bands of absorption are usually identified (Figure 3). The G band (~1580 cm<sup>-1</sup>) is typically assigned to a graphitic lattice mode E<sub>2g</sub> [35,36].

Figure 2. Representative micro-Raman spectra of the investigated macerals.
Figure 3. Curve-fitting of a representative attrinite spectrum.

However, in the case of lignite, it is mainly related to aromatic ring breathing (Li et al., 2006). The D2 band (~1615 cm$^{-1}$) makes a shoulder on the G band, being also attributed to the graphitic E$_{2g}$ mode [37,38]. It always occurs when the D1 band is present [39]. The D3 band (~1540 cm$^{-1}$), also referred to as G$_{R}$ (Li et al., 2006), originates from out-of-plane vibration due to structural defects and heteroatoms [40,41]. It is assigned to aromatic systems (composed of three to five rings) and amorphous carbon [6,37,42,43].

The D6 band (~1465 cm$^{-1}$), also denoted as V$_{L}$ [6], is related to methyl or methylene, small aromatic systems, and amorphous carbon [6]. The D1 band (~1285 cm$^{-1}$) is assigned to in-plane defects or occurrence of heteroatoms and is associated with the breathing mode (A$_{1g}$) of sp$^{2}$ atoms in aromatic rings [35,38,41]. The origin of the D5 band (~1285 cm$^{-1}$) or S$_{L}$ band [6] is not fully understood. It is believed to represent aryl–alkyl ethers [6] or CH in aliphatic chains [44]. The D4 band (~1195 cm$^{-1}$) (S band—in Li et al., 2006) is attributed mainly to the occurrence of sp$^{3}$-rich or sp$^{3}$–sp$^{2}$ carbon structures, alkyl–aryl groups, aliphatic (or aromatic) ethers, as well as CH in aliphatic chains or on aromatic rings [6,43–45]. Frequently, especially in bituminous coals and cokes, only one band in the 1200–1240 cm$^{-1}$ region, named D4, is indicated [43,46–49]. A very weak D7 band at ~1095 cm$^{-1}$, marked S$_{R}$ by Li et al. [6], is found in most of the spectra. It is assigned to C-H on aromatic rings [6]. Additionally, a very weak band at ~1700 cm$^{-1}$, related to the
carbonyl group (C=O) vibration, was also observed in some of the spectra (not used in the curve-fitting procedure).

Similar Raman bands were observed in the earlier Raman investigations of low-rank coal (ignite and sub-bituminous coal) [6–8,10,11,44]. However, the $V_R$ band at ~1380 cm$^{-1}$ indicated by Li et al. [6] and Xu et al. [11] was not found.

The G band in the attrinite spectra appears at ~1579 cm$^{-1}$, whereas the D1 band appears at ~1376 cm$^{-1}$ (Table 4). The full width at half maximum (FWHM) of these bands is ~79 cm$^{-1}$ and ~138 cm$^{-1}$, respectively. The Raman Band Separation (RBS) value is ~203 cm$^{-1}$ (Table 5).

In the case of densinite, the G band is found at ~1580 cm$^{-1}$, and the D1 band at ~1374 cm$^{-1}$. The FWHM of these bands is ~78 cm$^{-1}$ and ~140 cm$^{-1}$, respectively (Table 4). The RBS reaches a value of ~206 cm$^{-1}$ (Table 5).

The G band in the ulminite spectra occurs at ~1581 cm$^{-1}$, while the D1 band occurs at ~1369 cm$^{-1}$, with FWHM values of ~80 cm$^{-1}$ and ~147 cm$^{-1}$, respectively (Table 4). The RBS value is ~213 cm$^{-1}$ (Table 5).

Considering fusinite spectra, the G band position is ~1584 cm$^{-1}$, whereas the D1 band position is ~1367 cm$^{-1}$; the FWHM of these bands is ~73 cm$^{-1}$ and ~151 cm$^{-1}$, respectively (Table 4). The RBS reaches ~217 cm$^{-1}$ (Table 5). The G band FWHM is higher and the RBS lower than those determined for fusinite and semifusinite in bituminous (hard) coals [47,50]. Fusinite spectra have a weak or absent D7 band at ~1095 cm$^{-1}$.

The values of other spectral parameters are summarized in Tables 4 and 5.

The sum of the $A_{D4}/A_{ALL} + A_{D5}/A_{ALL}$ ratio calculated herein is similar to the value of the $A_{D4}/A_{ALL}$ ratio (derived from a typical five-band curve fitting method) obtained for semifusinite, but lower than that of fusinite in bituminous coals [47,50]. Furthermore, the sum of the $A_{D4}/A_{ALL} + A_{D5}/A_{ALL}$ ratios determined in this study corresponds well to the value of the $A_{D3}/A_{ALL}$ ratio (one band was indicated in the valley between the D1 and G bands) in the spectra of semifusinite from bituminous coals [47,50].

### Table 4. Band positions and full width at half maximum (FWHM) in the spectra of the studied macerals.

| Maceral      | D2 pos. cm$^{-1}$ | D2 FWHM cm$^{-1}$ | G pos. cm$^{-1}$ | G FWHM cm$^{-1}$ | D3 pos. cm$^{-1}$ | D3 FWHM cm$^{-1}$ | D6 pos. cm$^{-1}$ | D6 FWHM cm$^{-1}$ | D1 pos. cm$^{-1}$ | D1 FWHM cm$^{-1}$ | D5 pos. cm$^{-1}$ | D5 FWHM cm$^{-1}$ | D4 pos. cm$^{-1}$ | D4 FWHM cm$^{-1}$ | D7 pos. cm$^{-1}$ | D7 FWHM cm$^{-1}$ |
|--------------|-------------------|-------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| Attrinite    | 1613.3            | 76.3              | 1578.7           | 78.7             | 1541.1           | 93.3             | 1471.5           | 127.0            | 1375.9           | 138.0            | 1297.7           | 137.5            | 1191.9           | 105.6            | 1101.7           | 74.7             |
| n = 28       | 2.3               | 5.0               | 2.6              | 4.6              | 10.6             | 6.6              | 13.8             | 2.9              | 14.1             | 4.8              | 18.2             | 4.2              | 15.9             | 6.3              | 25.00            |
| Densinite    | 1614.6            | 74.0              | 1579.7           | 77.8             | 1541.6           | 95.8             | 1468.1           | 139.7            | 1374.1           | 140.7            | 1289.9           | 138.0            | 1197.1           | 126.9            | 1098.8           | 105.0            |
| n = 30       | 2.4               | 4.3               | 2.9              | 4.0              | 3.9              | 10.5             | 8.5              | 13.5             | 3.2              | 16.3             | 5.0              | 20.5             | 7.3              | 37.9             | 8.3              | 77.3             |
| Ulminite     | 1616.8            | 73.6              | 1581.5           | 79.7             | 1539.2           | 103.3            | 1461.4           | 142.1            | 1366.5           | 146.8            | 1299.4           | 131.9            | 1199.0           | 134.6            | 1094.6           | 87.1             |
| n = 28       | 3.0               | 5.2               | 2.7              | 4.5              | 3.1              | 6.9              | 7.00             | 14.4             | 4.2              | 17.3             | 5.8              | 17.1             | 7.6              | 34.5             | 11.1             | 37.8             |
| Fusinite     | 1612.0            | 52.9              | 1583.7           | 72.8             | 1538.4           | 123.8            | 1458.4           | 132.8            | 1366.6           | 151.4            | 1278.5           | 153.6            | 1189.2           | 124.5            | 1088.4           | 108.0            |
| n = 27       | 2.2               | 6.5               | 2.7              | 4.8              | 5.7              | 15.4             | 10.9             | 26.7             | 3.3              | 10.6             | 8.7              | 23.8             | 9.3              | 31.8             | 8.8              | 29.5             |

Explanation: $n$—number of measurements; the standard deviation of the measurements is given in italics.

### Table 5. Spectral ratios and Raman Band Separation (RBS) derived from the curve fitting procedure.

| Maceral      | $I_{D1}/I_G$ | $A_{D3+D6}/A_{ALL}$ | $A_{D5}/A_{ALL}$ | $A_{D4}/A_{ALL}$ | RBS cm$^{-1}$ |
|--------------|-------------|---------------------|------------------|------------------|---------------|
| Attrinite    | 1.05        | 0.71                | 0.18             | 0.05             | 202.8         |
| n = 28       | 0.08        | 0.03                | 0.04             | 0.01             | 4.2           |
| Densinite    | 1.04        | 0.73                | 0.16             | 0.06             | 205.6         |
| n = 30       | 0.14        | 0.03                | 0.04             | 0.02             | 4.1           |
| Ulminite     | 0.95        | 0.27                | 0.14             | 0.07             | 212.9         |
| n = 28       | 0.12        | 0.04                | 0.04             | 0.03             | 5.4           |
| Fusinite     | 1.09        | 0.15                | 0.15             | 0.05             | 217.0         |
| n = 27       | 0.16        | 0.03                | 0.04             | 0.02             | 6.6           |

Explanation: $n$—number of measurements; the standard deviation of the measurements is given in italics.
4.3.2. Comparison of the Microstructural and Chemical Properties of the Analyzed Macerals

Comparisons were performed for the following pairs of macerals: attrinite–densinite, attrinite–ulminite, and attrinite–fusinite. To find out if the differences in spectral parameters between the macerals were statistically valid, the Shapiro–Wilk, Fisher–Snedecor, and, finally, Student’s t-test were performed.

The Shapiro–Wilk test shows that all data sets of individual Raman-derived spectral parameters obtained from the curve fitting procedure have normal or close to normal distribution. The Fisher–Snedecor test demonstrates that variances are equal in all analyzed pairs of spectral parameters.

Considering this, Student’s t-test was conducted for all parameters. Each time that the difference in spectral parameters is mentioned in the text, it is statistically significant, as inferred from the t-test (p value < 0.05).

Attrinite and densinite show the highest similarity of microstructural properties among all of the analyzed macerals, which is in line with the results of the FTIR investigations revealing that the chemical composition of detrohuminite subgroup macerals is largely independent of its degree of gelification [51]. However, there are some statistically significant (p < 0.05) differences between their spectral parameters. This applies to the position of the D1 band, which appears at lower wavenumbers in the densinite spectra, whereas the D4 band falls at higher wavenumbers (Figure 4a,b). The densinite spectra also reveal higher D6 and D4 band FWHMs and higher $A_{D3+D6}/A_{ALL}$ and $A_{D4}/A_{ALL}$ ratios (Figure 4c–f). This reflects higher amounts of small aromatic units and alkyl–aryl and CH groups in the chemical structure of densinite compared with attrinite [6,43–45], probably resulting from stronger gelification of densinite. An increase in aromaticity of the huminite group macerals with increasing gelification was earlier reported, based on the FTIR examination [5,52–54]. This is also in line with the difference in $R_r$ value between densinite (0.31%) and attrinite (0.26%). Similarly, the difference in D1 band position observed herein corresponds with a shift of the D1 band to lower wavelengths with increasing reflectance, as reported by Kelemen and Fang [55] and Guedes et al. [8]. The RBS value, which correlates well with the $R_r$ value [32,55–59], is higher for densinite, which also corresponds with higher reflectance values of this maceral (Tables 2 and 5). Where detrohuminite’s microstructure is concerned, it should be remembered that the primary plant material and the initial peat-forming conditions have a great impact on the features of macerals belonging to this subgroup.

Statistical analysis shows that the microstructural features of attrinite and ulminite differ significantly. The position of some of the Raman bands (D2, G, and D4) in ulminite spectra is moved to higher wavenumbers, whereas the D6, D1, and D7 bands fall at lower ones (Figure 5a,b; Table 4). Ulminite exhibits a higher FWHM of the D3, D6, D1, and D4 bands and higher $A_{D3+D6}/A_{ALL}$ and $A_{D4}/A_{ALL}$ ratios in comparison with attrinite (Figure 5c–e; Tables 4 and 5). On the other hand, the $A_{D5}/A_{ALL}$ ratio is lower in ulminite than in attrinite (Figure 5f). Taking the origin of the D3, D4, and D6 bands [6,43–45], the differences in the values of the spectral ratios imply that the chemical structure of ulminite is more abundant in small aromatic systems composed of three to five rings, C_aromatic–C_alkyl systems, and CH groups. This may be due to gelification of ulminite, as aromacity increases with increasing gelification [5,52–54]. Ulminite spectra also exhibit higher Raman band separation (RBS) values (Table 5) than those of attrinite. The differences observed may also result from different origins of starting materials; attrinite may come from the soft, cellulose-rich tissues of plants, and ulminite may come from lignin-rich xylem. Lignin remnants may affect the aromacity of lignite at the early stage of coalification [60].
Figure 4. Comparison of Raman spectral parameters obtained for attrinite and densinite: D1 band position (a), the D4 band position (b), the D6 band FWHM (c), the D4 band FWHM (d) the AD3+D6/AALL ratio (e) and AD4/AALL ratio (f). Explanation: pos.—position; A—attrinite; D—densinite.
Figure 5. Comparison of Raman spectral parameters obtained for attrinite and ulminite: G band position (a), the D1 band position (b), the D1 band FWHM (c), the \( \frac{A_{D3+D6}}{A_{ALL}} \) ratio (d), \( \frac{A_{D4}}{A_{ALL}} \) ratio (e) and the \( \frac{A_{D5}}{A_{ALL}} \) ratio (f). Explanation: pos.—position; A—attrinite; U—ulminite.

As should be expected, there are major differences in microstructure between attrinite and fusinite. Most of the Raman bands occur at lower wavenumbers when fusinite is considered, but the position of the G band moves to higher wavenumbers (Figure 6a,b; Table 4). Fusinite spectra also have lower FWHM values of the D2 and G bands and a higher FWHM of the D3, D1, D5, and D7 bands in comparison with attrinite (Figure 6c,d; Table 4). However, the D3, D6, and D5 bands reveal low height; thereby, the \( \frac{A_{D3+D6}}{A_{ALL}} \) and \( \frac{A_{D5}}{A_{ALL}} \) ratios are lower for fusinite (Figure 6e,f; Table 5). The RBS value determined for fusinite is also higher (Table 5). The lower G band FWHM indicates better organization of
the fusinite microstructure in relation to the attrinite microstructure [8,32,39,47,55,61,62]. The higher D1 band FWHM in the fusinite spectra reveals that aromatic clusters in fusinite are more varied in terms of their order and dimensions [63]. The higher RBS value determined for fusinite corresponds with the higher mean random reflectance (Rr) of this maceral (Tables 2 and 5) [32,55–59]. The lower \( \frac{A_{D3}}{A_{ALL}} \), \( \frac{A_{D6}}{A_{ALL}} \), and \( \frac{A_{D5}}{A_{ALL}} \) ratio values determined for fusinite (when compared with attrinite) are a consequence of the lower content of smaller aromatic systems—aliphatics as well as mixed structures and alkyl–aryl groups [6,37,42–44]. Instead, larger aromatic ring systems dominate the chemical structure of fusinite.

**Figure 6.** Comparison of Raman spectral parameters obtained for attrinite and fusinite: G band position (a), the D1 band position (b), the G band FWHM (c), the D1 band FWHM (d), the \( \frac{A_{D3+D6}}{A_{ALL}} \) ratio (e), and the \( \frac{A_{D5}}{A_{ALL}} \) ratio (f). Explanation: pos.—position; A—attrinite; F—fusinite.
Ferralis et al. [44] found a linear relationship between the \((I_{D4}+I_{D5})/(I_G+I_{D2})\) Raman band area ratio (Symbols taken from Ferralis et al. [44]) and the elemental H:C ratio. Considering the above, a similar calculation of the \((I_{D4}+I_{D5})/(I_G+I_{D2})\) ratio was performed here, which shows much lower values obtained for fusinite (0.26) than for attrinite (0.37), densinite (0.46), or ulminite (0.50). This reflects the much lower hydrogen content in the chemical structure of fusinite in comparison with other examined macerals. However, no positive correlation between the \(A_{D4}/A_{ALL}\) and \(A_{D5}/A_{ALL}\) ratios is observed—which should be expected if the D4 and D5 bands are related to the same chemical structures in coal. This may suggest that the origin of the D4 and D5 bands is different.

5. Conclusions
Lignite (ulminite reflectance \(R_{f0} = 0.27\%\)) from the Szczerców deposit (Central Poland) was subjected to micro-Raman spectroscopy examination. The study revealed differences in the microstructure of the huminite group macerals (attrinite, densinite, and ulminite). Densinite and ulminite are more abundant in small aromatic units than attrinite, which may result from their gelification or the kind of starting material. The differences in the Raman spectral characteristics between attrinite and ulminite are more pronounced than between attrinite and densinite. As expected, fusinite, in comparison with the huminite group macerals, is composed of larger, more varied aromatic systems. Similarities to semifusinite’s microstructure in bituminous coals are observed. The D4 band (1190–1200 cm\(^{-1}\)) and D5 band (1280–1290 cm\(^{-1}\)) most likely correspond to different chemical structures, and their origin should be further investigated.

On the basis of petrographic analysis, the directions of use and the best ways of exploitation can also be determined. The presence of fibrous xylites is particularly unfavorable during mining and grinding. Therefore, in practice, the best solution is to determine the xylite content as early as possible, even at the stage of deposit exploration. The petrographic composition of coal has a direct impact on its suitability for the gasification process, where the degree of coal gelification and the concentration of carbonate and quartz minerals are of particular importance.

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Appendix A

Figure A1. Selected measurement points (+) on attrinite (A–F).
Figure A2. Selected measurement points (+) on densinite (A–F).
Figure A3. Selected measurement points (+) on ulminite (A–F).
Figure A4. Selected measurement points (+) on fusinite (A–F).

References
1. Suárez-Ruiz, I.; Crelling, J. Applied Coal Petrology; Elsevier Ltd.: Amsterdam, The Netherlands, 2008; ISBN 9780080450513.
2. Suárez-Ruiz, I.; Rubiera, F.; Diez, M. New Trends in Coal Conversion: Combustion, Gasification, Emissions, and Coking; Woodhead Publishing: Cambridge, UK, 2018.
3. Roberts, M.J.; Everson, R.C.; Neomagus, H.W.J.P.; Van Niekerk, D.; Mathews, J.P.; Branken, D.J. Influence of maceral composition on the structure, properties and behaviour of chars derived from South African coals. Fuel 2015, 142, 9–20. [CrossRef]
4. Bielowicz, B.; Baran, P. CO₂ sorption properties of selected lithotypes of lignite from Polish deposits. Geol. Q. 2019, 63, 786–800. [CrossRef]
5. Wagner, M. The character of the IR absorption in the spectral rang 1700–1500 cm⁻¹ of some macerals of the huminite group of brown coal. Bull. L’Acad. Pol. Sci. 1981, 29, 321–330.
6. Li, X.; Hamashi, J.I.; Li, C.Z.; Hayaschi, J.; Li, C.Z. FT-Raman spectroscopic study of the evolution of char structure during the pyrolysis of a Victorian brown coal. Fuel 2006, 85, 1700–1707. [CrossRef]

7. Liu, X.; You, J.; Wang, Y.; Lu, L.; Xie, Y.; Yu, I.; Fu, Q. Raman spectroscopic study on the pyrolysis of Australian bituminous coal. J. Fuel Chem. Technol. 2014, 42, 270–276. [CrossRef]

8. Guedes, A.; Valentim, B.; Prieto, A.C.; Rodrigues, S.; Noronha, F. Micro-Raman spectroscopy of colliotelinite, fusinite and macrinite. Int. J. Coal Geol. 2010, 83, 415–422. [CrossRef]

9. Bayyal; M.; Yüreu̇, A.; Yildiz, B.; Yüreu̇, Y. Structure of some western Anatolia coals investigated by FTIR, Raman, 13C solid state NMR spectroscopy and X-ray diffraction. Int. J. Coal Geol. 2016, 165, 166–176. [CrossRef]

10. He, X.; Liu, X.; Nie, B.; Song, D. FTIR and Raman spectroscopy characterization of functional groups in various rank coals. Fuel 2017, 206, 555–563. [CrossRef]

11. Xu, J.; Tang, H.; Su, S.; Liu, J.; Xu, K.; Qian, K.; Wang, Y.; Zhou, Y.; Hu, S.; Zhang, A.; et al. A study of the relationships between coal structures and combustion characteristics: The insights from micro-Raman spectroscopy based on 32 kinds of Chinese coals. Appl. Energy 2018, 212, 46–56. [CrossRef]

12. Kasinski, J.; Piwocki, M.; Sadowska, E.; Ziemiańska-Tworydło, M. Lignite of the Polish Lowlands Miocene: Characteristics on a base of selected profiles. Biul. Państw. Inst. Geol. 2009, 1, 3–22. [CrossRef]

13. Wagner, M. Zmienność Petrologiczno-Sedymentologiczna i Własności Technologiczne Kredy Jeziornej w Osadach Neogenu Typu Wapiennego Zapadliska Tecktonicznego na Przykładzie Złoża Węgla Brunatnego “Szczerców” (Petological and Sedimentological Variability and Tech; AGH Uczelniane Wydawn. Naukowo-Dydaktyczne: Kraków, Poland, 2007; ISBN 9788374641289.

14. Wagner, M.; Matl, K. Stratygrafia Kredy Jeziornej ze Złoża Węgla Brunatnego “Szczerców”. (Stratigraphy of Lacustrine Chalk from the “Szczerców” Lignite Deposit); Wydawnictwa AGH: Kraków, Poland, 2007; Volume 33.

15. Piwocki, M.; Ziemiańska-Tworydło, M. Neogene of the polish lowlands—Lithostratigraphy and pollen-spore zones. Kwart. Geol. 1997, 41, 21–40.

16. Widera, M. An overview of lithotype associations of Miocene lignite seams exploited in Poland. Geologos 2016, 22, 213–225. [CrossRef]

17. Słomka, T.; Wagner, M. Charakter Petrograficzny I Warunki Sedymentacji Wybranych Kompleksów Litologicznych Z Profili Miocenu W złożu węGla Brunatnego Belchatów (Petrological Studies and Sedimentological Conditions of Select Lithologic Series from Miocene from Belchatów W Lign. Prace Geol. Pan 2000, 147. Available online: https://integro.cniba.edu.pl/integro/191800753249/ksiazka/charakter-petrograficzny-i-warunki-sedymentacji-wybranych-kompleksow-litologicznych-z-profilu-miocenu?autor=ksika&ln=gb&_lang=en (accessed on 24 December 2020).

18. Kasinski, J.R.; Piwocki, M. Low-rank coals in Poland: Prospection—Mining—Progress. Pol. Geol. Inst. Spec. Pap. 2002, 7, 17–30.

19. ISO 7404-2. Methods for the Petrographic Analysis of Coals—Part 2: Methods of Preparing Coal Samples; ISO: Geneva, Switzerland, 2009.

20. ISO 7404-3. Methods for the Petrographic Analysis of Coals—Part 3: Method of Determining Microscopically the Reflectance of Vitrite; ISO: Geneva, Switzerland, 2009.

21. ICCP. The new inertinite classification (ICCP System 1994). Fuel 2001, 80, 459–471. [CrossRef]

22. Pickel, W.; Kus, J.; Flores, D.; Kalaitzidis, S.; Christian, K.; Cardott, B.J.; Misz-Kennan, M.; Rodrigues, S.; Hentschel, A.; Hamor-Vido, M.; et al. Classification of liptinite—ICCP System 1994. Int. J. Coal Geol. 2017, 169, 40–61. [CrossRef]

23. Sykorová, I.; Pickel, W.; Christian, K.; Wolf, M.; Taylor, G.H.; Flores, D. Classification of huminite—ICCP System 1994. Int. J. Coal Geol. 2005, 62, 85–106. [CrossRef]

24. ISO 7404-3. Methods for the Petrographic Analysis of Coals—Part 3: Method of Determining Maceral Group Composition; ISO: Geneva, Switzerland, 2009.

25. ISO 579. Coke—Determination of Total Moisture; ISO: Geneva, Switzerland, 2013.

26. ISO 11760. Classification of Coals; ISO: Geneva, Switzerland, 2005.

27. UN ECE. International Classification of in-Seam Coals; UN ECE: Geneva, Switzerland, 1998.

28. Tuinstra, F.; Koenig, J.L. Raman Spectrum of Graphite. J. Chem. Phys. 1970, 53, 1126–1130. [CrossRef]

29. Nemanich, R.J.; Solin, S.A. First- and second-order Raman scattering from finite-size crystals of graphite. Phys. Rev. B 1979, 20, 392–401. [CrossRef]

30. Cuesta, A.; Dhamelincourt, P.; Laureyns, J.; Martinez-Alonso, A.; Tascon, J.M.D. Raman microprobe studies on carbon materials. Carbon 1994, 32, 1523–1532. [CrossRef]
