

Originally published as:

Nadoll, P., Rehm, M., Duschl, F., Klemd, R., Kraemer, D., Sosnicka, M. (2018): REY and Trace Element Chemistry of Fluorite from Post-Variscan Hydrothermal Veins in Paleozoic Units of the North German Basin. - *Geosciences*, *8*, 8.

DOI: http://doi.org/10.3390/geosciences8080283



Article

# **REY and Trace Element Chemistry of Fluorite from Post-Variscan Hydrothermal Veins in Paleozoic Units of the North German Basin**

Patrick Nadoll <sup>1,\*</sup><sup>(D)</sup>, Meike Rehm <sup>1</sup>, Florian Duschl <sup>2</sup>, Reiner Klemd <sup>1</sup>, Dennis Kraemer <sup>3</sup><sup>(D)</sup> and Marta Sośnicka <sup>4</sup>

- <sup>1</sup> GeoZentrum Nordbayern, Friedrich-Alexander-University Erlangen-Nuremberg, Schlossgarten 5, 91054 Erlangen, Germany; meike.rehm@web.de (M.R.); reinerklemd@fau.de (R.K.)
- <sup>2</sup> Geowissenschaftliches Zentrum der Universität Göttingen, Goldschmidtstrasse 1-3, 37077 Göttingen, Germany; florian.duschl@geo.uni-goettingen.de
- <sup>3</sup> Department of Physics and Earth Sciences, Jacobs University Bremen, Campus Ring 1, 28759 Bremen, Germany; d.kraemer@jacobs-university.de
- <sup>4</sup> GFZ, German Research Centre for Geosciences, Telegrafenberg, 14473 Potsdam, Germany; sosnicka@gfz-potsdam.de
- \* Correspondence: pnadoll@gmail.com

Received: 19 June 2018; Accepted: 26 July 2018; Published: 29 July 2018



Abstract: Hydrothermal fluorites from Paleozoic sedimentary rocks and volcanic units in the North German Basin (NGB) have been investigated to create a petrographic and geochemical inventory-with particular focus on strategic elements such as rare earth elements (REE)-and to uncover possible links between the post-Variscan hydrothermal mineralization in the NGB and bordering areas such as the Harz Mountains and Flechtingen Calvörde Block (FCB). Fluorites from ten localities underwent a detailed petrographic examination, including SEM-BSE/CL imagery, and were compositionally analysed using LA-ICP-MS. Overall, REY concentrations are comparatively low in fluorite from all investigated areas—the median sum of REY ranges from 0.3 to 176 ppm. Europium<sub>CN</sub> anomalies are slightly negative or absent, indicating that either the formation fluid experienced temperatures above 250 °C or that fluid-rock interactions and REE enrichment was likely controlled by the source rock (i.e., volcanic) composition and complexation processes. Fluorites from the Altmark-Brandenburg Basin (ABB) and the Lower Saxony Basin (LSB) display distinctly different REY<sub>CN</sub> signatures, suggesting that fluid compositions and genetic processes such as fluid-rock interaction differed significantly between the two areas. Complex growth zones and REY<sub>CN</sub> signatures in fluorite from the ABB and the FCB reflect geochemical variability due to adsorption processes and intrinsic crystallographic controls and imply that they are genetically related. Two petrographically and geochemically distinct generations are observed: Fluorite I—light SEM shades, relatively enriched in LREE; Fluorite II—darker SEM shades, comparatively depleted LREE, slightly higher HREE concentrations. These fluorite generations represent zoned (or cyclical) growth within a single progressive hydrothermal event and do not reflect a secondary remobilization process. We demonstrate that increasing Tb/La ratios and decreasing La/Ho ratios can be the result of continuous zoned growth during a single mineralizing event, with significant compositional variations on a micron-scale. This has implications for the interpretation of such trends and hence the inferred genetic evolution of fluorite that displays such geochemical patterns. The complex micro-scale intergrowth of these generations stresses the need for detailed petrographic investigations when geochemical data are collected and interpreted for mineral exploration.

**Keywords:** fluorite; hydrothermal veins; North German Basin; rare earth elements; trace elements; zoned growth; remobilization



#### 1. Introduction

As part of the r4 national collaborative research effort that explores the resource potential of strategic elements such as rare earth elements (REE) in Germany, we investigated hydrothermal fluorite mineralization in Paleozoic sedimentary rocks and volcanic units of the North German Basin (NGB). Although the structure, stratigraphy and evolution of the NGB and its sub-basins are well-studied due to extensive exploration and production drilling programs, hydrothermal mineralization, including fluorite-barite and base metal sulphide mineralization in different parts of the basin, have not been studied in detail until recently [1-3]. Here, we present macro- and micro-scale petrographic observations (optical microscopy, back-scattered secondary electron (BSE) and cathodoluminescence (CL) imaging) in combination with geochemical data (laser ablation inductively coupled mass spectrometry (LA-ICP-MS), bulk rock X-ray fluorescence (XRF)), including REE and minor and trace element contents, for hydrothermal fluorite from four distinct areas within the NGB—specifically, the Altmark-Brandenburg Basin (ABB) as part of the North East German Basin (NEGB), the Lower Saxony Basin (LSB) in the western part of the NGB, the Flechtingen High (Flechtingen Calvörde Block, FCB) and the Harz Mountains south of the ABB. The latter two were selected to investigate possible links between the hydrothermal mineralization in the NGB and the surrounding areas. In addition to the characterization of the fluorite mineralization and the observed characteristic geochemical signatures, we present and discuss the micro-scale fluorite mineralogy and the corresponding elemental distribution which can have considerable impact on the interpretation of bulk and in-situ fluorite data regarding the evolution of the hydrothermal system.

#### 1.1. General Setting

The NGB is a post-Variscan intracratonic structure formed by crustal thinning accompanied by extensive volcanic activity (Upper Carboniferous to Lower Permian) and later inversion (Cretaceous) which is also evident in the uplift of the FCB and Harz Mountains south of the NEGB (e.g., [4–10]). Changing tectonic settings, from extensional to divergent movements and the stepwise and long-term subsidence followed by a later inversion of the basin enabled fluid migration along structurally controlled pathways and led to the formation of several hydrothermal vein systems in the NGB [2,11–15]. The possible fluid sources (e.g., meteoric or magmatic water) and the heat source for the enhanced geothermal gradients are also discussed by these authors. Extensive fault systems provided pathways for migrating fluids and convective fluid flow in Rotliegend (Lower Permian) and Upper Carboniferous units [1,11,15,16] which was further enhanced during basin inversion, also allowing for basinal brines of various salinities to mix e.g., [16]. In particular the fluorite-barite mineralization and REE-enriched hematite veins in the ABB and FCB have been attributed to the latter phase [13,17]. The Harz Mountains have historically been an important mining area for fluorite and other commodities and post-Variscan hydrothermal mineralization in this area shows many similarities to the hydrothermal veins found in sedimentary rocks and volcanic units further north in the FCB and the ABB [12,18]. For example, sulphur isotopic ratios measured by Schmidt Mumm and Wolfgramm [13] are strikingly similar and could reflect a mutual source, that is, Rotliegend volcanics and underlying Carboniferous sedimentary rocks. Hence, we compare the hydrothermal fluorite from the ABB, FCB and LSB with a quartz-calcite-fluorite vein from the Biwender Vein Complex near Straßberg in the Harz Mountains. This area has a mining history that dates back to the 1300s, with fluorite, lead and silver being the main resources [19].

## 1.2. Fluorite (Ca<sub>2</sub>F) Geochemistry

Fluorite is a common accessory and gangue mineral in many magmatic and hydrothermal deposits and can also occur as a primary economic mineral in massive or vein-type mineralization (e.g., Vergenoeg [20], Pennine Orefield [21,22] and Wölsendorf [23]). It has been the subject of many studies investigating its unique geochemical signature as a function of its formation history

e.g., [24–33]. Many of these investigations focused on the REE + Y (REY) concentrations in fluorite because the behaviour of these elements follows well established trends and is controlled by a number of known variables such as temperature, fluid composition, fluid-rock interaction and intrinsic crystallographic controls [23,32,34–40]. Fluorites and other Ca-bearing minerals such as calcite may preserve the REY signature of the hydrothermal fluid and hence the REY signature of the source [36]. REY are largely unaffected by water-rock interactions and are therefore independent of the host rock composition [29,30,36]. This makes them a valuable proxy for the formation conditions and the initial fluid composition [29]. This information may include the metal source, temperature conditions of water-rock interactions, migration, precipitation and the chemical composition of the fluid phase [29,32,33,36,41]. Möller [33] introduced the Tb/Ca-Tb/La diagram which can discriminate between sedimentary, hydrothermal and pegmatitic fluorite and allows the differentiation between primary and recrystallized fluorite [32,38,42]. Further relevant examples are studies by Bau et al. [32] and Kraemer et al. [43] who used REY patterns and Sr-Nd-Pb isotopes in fluorites from the Pennine MVT ore field to demonstrate that temperature and sources of the fluorite-forming fluids differed significantly between the Southern Pennine Orefield and the Northern Pennine Orefield. Fluorites in the Alston Block of the Northern Pennine Orefield are about ten-fold enriched in REY (553 mg kg<sup>-1</sup>; [43]) relative to fluorites from the Askrigg Block in the Northern Pennine Orefield (42.7 mg kg<sup>-1</sup>, [43]) and fluorites from the Southern Pennine Orefield (28 mg kg<sup>-1</sup>; [32]).

In addition to REY, the Ca<sup>2+</sup> cation in the fluorite crystal structure can be substituted by a range of other foreign cations including Li<sup>+</sup>, Na<sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, Mn<sup>2+</sup>, Fe<sup>2+,3+</sup>, Zn<sup>2+</sup>, Sr<sup>2+</sup>, Y<sup>3+</sup>, Zr<sup>4+</sup>, Ba<sup>2+</sup>, Pb<sup>2+</sup>, Th<sup>4+</sup>, and U<sup>4+</sup> e.g., [29,36,44–46]. The overall concentrations of these cations usually occur at minor to trace element levels, that is, they do not commonly exceed 1 wt.% [47].

## 2. Geological Setting

The three main study areas, that is, the Altmark-Brandenburg Basin (ABB), the Flechtingen Calvörde Block (FCB) and the Lower Saxony Basin (LSB) are briefly introduced (see also Figure 1).

## 2.1. North German Basin

The North German Basin (NGB) is part of the supra-regional Central European Basin System, which developed after the break-down of Pangea [9,48]. The German part is formed by the Variscan Foreland Basin of Late Carboniferous age, including the NGB, the LSB of Late Jurassic to Early Cretaceous age and the North Sea Basin. Rotliegend sandstones and Zechstein (Upper Permian) carbonates within the NGB host hydrocarbon accumulations [49] and have been drilled extensively by the oil and gas industry. A number of these drill cores feature hydrothermal mineralization, including base metal sulphide and fluorite-barite veins, in deeply buried Paleozoic sedimentary rocks of the NGB (ca. 3–4 km), specifically the ABB and the LSB.

# 2.2. Altmark-Brandenburg Basin

The Flechtinger Fault Zone and the Gifhorn Trough mark the southern and western border of the ABB, respectively [50,51] (Figure 1). The ABB is commonly considered devoid of any known economically viable mineralization. However, hydrothermal vein-type F-Ba mineralization hosted by Rotliegend volcanic rocks [7] and Permian siliciclastic sedimentary rocks were found in gas exploration drill cores in the Altmark area e.g., [52].



**Figure 1.** Location of sampled wells/localities and major structural features of northern Germany, including the ABB, the LSB, the FCB (Flechtingen High) and the Harz. The thin unlabelled dashed line indicates the extend of the NGB with its shallow platform area (dashed area). Well names are anonymized, and locations are approximate.

#### 2.3. Lower Saxony Basin

The E–W trending LSB is located in the NW-part of Germany (Figure 1), north of the Rhenish Massif [5,53]. The LSB developed during a tectonically active period in Late Jurassic to Mid-Cretaceous times along NW–SE trending fault systems, superimposing the Late Carboniferous Variscan Foreland Basin and parts of the Variscan Fold Belt [5,54,55]. During the Cretaceous uplift and inversion, the LSB was in part overthrust on the southern Muensterland Block along the Osning Thrust Zone [56–58]. Faults of Permian age limit the Osning Thrust Zone to the south [59].

# 2.4. Flechtingen Calvörde Block and Harz Mountains

The Harz Mountains and the FCB are expressions of the Variscan Fold Belt within the Rhenohercynian Zone e.g., [18,60] (Figure 1). The FCB to the northeast is divided from the Harz by the Subhercynian Basin [18]. The Harz Mountains are located along the southern margin of the EFZ, between the two major sub-basins, the LSB in the northwest and the ABB in the northeast (Figure 1). The occurrence of major fault-related hydrothermal mineralization in the Harz Mountains such as the Biwender Vein and their association with the complex post-Variscan tectonic history may be connected with fluid migration in the adjacent sedimentary NGB [61,62]. The Harz Mountains were uplifted during the Late Cretaceous-Early Cenozoic contemporaneously with the NGB inversion e.g., [63,64].

The formation of vein mineralization in the Harz Mountains developed from the Late Variscan through to Upper Cretaceous times. Hydrothermal vein-type Pb-Zn (Ag), Ba-F, Ba-Fe and F-Ba mineralization was deposited in the Mesozoic (Triassic through Late Cretaceous) e.g., [65–67]. Post-Variscan, Late Triassic-Jurassic sulphide mineralization was preceded by intense hematitization and silicification (pre-ore stage) followed by the deposition of typically banded quartz and carbonate-sulphide veins [68]. Post-Variscan fluorite veins, such as the Biwender Vein which is hosted

in Carboniferous siliciclastic sedimentary rocks, have been attributed to the mixing of low-salinity, low-temperature formation waters with ascending high-salinity and high-temperature brines [42]. This area is also known for its historic Pb-Zn and Ag mining [19]. Hydrothermal Ba-F, F-Ba, and specular hematite vein-type mineralization within the FCB is hosted by Upper Carboniferous and lower Permian rhyolitic and andesitic volcanic rocks, which underwent intense alteration, that is, bleaching, oxidation, and sericitization [17]. These volcanic rocks host steeply-dipping hematite veins which are exposed in the Bodendorf quarry, 35 km northwest of Magdeburg. The length of these NW–SE-trending veins reaches several hundred meters and widths of up to 1.5 m [69]. Fluorite-bearing veins hosted by andesites most probably reflect hydrothermal activity induced by granite intrusion of uncertain (Permian?) age [70]. Schmidt Mumm & Wolfgramm [13,14] suggested a similarity between this hematite mineralization and iron oxide copper gold (IOCG) deposits formed in intracratonic basins. However, the character of the iron ore-forming mechanisms, the direct age of mineralization, and the iron source for the hematite veins remain unknown.

Outcrops of stratigraphic units of interest are scarce in our study areas, however, three quarries located in volcanic basement units of the FCB were investigated to compare fluid systems and mineralization ages as well as related fault systems with those found in drill cores from the adjacent ABB.

# 3. Methodology

Seventeen fluorite samples from the ABB and LSB (drill core samples) and the FCB and Harz Mountains (hand samples) have been investigated using the following array of analytical techniques. Polished mounts and thin sections of each sample were also investigated with transmitted and reflected light microscopy.

REY data are presented in C1 chondrite-normalized CN [71] spidergrams (REY<sub>CN</sub> plots).

# 3.1. SEM-BSE

The scanning electron microscope (SEM) was first and foremost used for the identification of small fluorite grains as well as the visualization of structural and textural features of fluorite. Thin sections and polished mounts were prepared with carbon coating and silver paint. For evaporation, the prepared samples were stored in a vacuum cabin for at least 24 h. The measurements were carried out at the GeoZentrum Nordbayern with a Tescan VEGA-II XMU coupled with a X-MAX EDS system from Oxford Instrument and a 50 mm<sup>2</sup> X-Max detector. Prior to analysis, quantum optimization with a Cu and Ti chip was carried out. Operating conditions are shown in Table 1 and also apply to the CL measurements.

Process Time	6 µs
Channels	2К
Energy spectrum	0–10 keV
Detector	BSE and CL mode
Accelerating voltage	15 kV
Working distance	13–15 mm 14.6 mm (optimum conditions)
Sample current	300–500 pA

Table 1. SEM operating conditions.

#### 3.2. Cathodoluminescence (CL)

CL was used to identify and illustrate internal structures and textures of fluorite grains. The wavelength in combination with the intensity of the emitted light can characterize individual minerals as well as the distribution of certain impurities within those minerals. Cathodoluminescence can be applied for distinguishing carbonate and quartz cement generations, and for contrasting microstructures related with fluid-rock interaction, like fluid pathways, paleo-porosity, diffusional

textures, healed microfractures [72,73]. These textures can normally not be observed by standard petrographical microscopy.

CL analyses were carried out on thin- and thick sections. At the GZG Göttingen a high-power HC3-LM- Simon-Neuser CL microscope [73] with a coupled Peltier-cooled Kappa PS 40C-285 (DX) camera system (resolution 1.5 mpx) attached to an Olympus BH-2 microscope was used. The electron gun was operated at a voltage of 14 keV under high vacuum ( $10^{-4}$  bar) with a filament current of 0.18 mA. The diameter of the electron beam was ca. 4 µm. Further CL measurements were carried out with the SEM at the GeoZentrum Nordbayern (see Table 1).

# 3.3. LA-ICP-MS

Laser ablation ICP-MS measurements were carried out on (1) bulk rock samples (fused discs) and (2) individual fluorite grains (polished mounts). The analyses were undertaken at the Bureau Veritas commercial laboratories in Perth and at the GeoZentrum Nordbayern.

- (1) For this investigation, a New Wave 193 μm eximer laser coupled to a 7700 agilent ICP-MS was used. The laser was equipped with a 2-volume cell. Each fused disc was ablated for 60 s with a laser spot-size of 150 μm in diameter, a pulse rate of 20 Hz, and measured fluence of 8 J/cm<sup>2</sup>. As carrier gas a mixture of He and Ar was used with a flow rate optimized for highest sensitivity. For calibration three reference materials were used. Two of these were manufactured and validated in-house at the Bureau Veritas laboratory, the third one was the OREAS 134b standard reference material.
- (2) An Agilent 7500c quadrupole ICP-MS coupled with a New Wave 193 µm excimer laser was used for analyses, with 35  $\mu$ m spot-size, a 16 Hz repetition rate and a fluence of 7.8 J/cm<sup>2</sup>. The background was measured for 20 s; the samples were analysed for 20 s. The NIST SRM 612 and NIST SRM 610 glasses were used as reference materials for fluorite. A stoichiometric value of 51.1 wt.% for Ca [74] was used as the internal standard for fluorite. Elemental concentrations, 1 sigma errors and minimum detection limits were calculated using GLITTER (Version 4.4.4, On-line Interactive Data Reduction for LA-ICPMS (Macquarie Research Ltd., Sydney, Australia) using the protocol of Griffin et al. [75]. For each element, the limit of quantification (LOQ) was set to three times the limit of detection (LOD). Signals lower than LOQ were not considered for further interpretation. The following isotopes were measured (typical LODs in ppm in brackets): <sup>23</sup>Na (1.28), <sup>29</sup>Si (43.8), <sup>42</sup>Ca, <sup>44</sup>Ca at 10 ms integration time each and <sup>7</sup>Li (0.11), <sup>24</sup>Mg (1.08), <sup>26</sup>Mg, <sup>39</sup>K (1.18), <sup>55</sup>Mn (0.13), <sup>57</sup>Fe (2.8), <sup>66</sup>Zn (0.41), <sup>71</sup>Ga (0.02), <sup>73</sup>Ge (0.08), <sup>74</sup>Ge, <sup>85</sup>Rb (0.005), <sup>88</sup>Sr (0.006), <sup>89</sup>Y (0.008), <sup>90</sup>Zr (0.017), <sup>93</sup>Nb (0.01), <sup>115</sup>In (0.008), <sup>118</sup>Sn (0.07), <sup>137</sup>Ba (0.06), <sup>139</sup>La (0.01), <sup>140</sup>Ce (0.01), <sup>141</sup>Pr (0.005), <sup>146</sup>Nd (0.03), <sup>147</sup>Sm (0.04), <sup>153</sup>Eu (0.005), <sup>157</sup>Gd (0.002), <sup>159</sup>Tb (0.003), <sup>163</sup>Dy (0.03), <sup>165</sup>Ho (0.004), <sup>166</sup>Er (0.01), <sup>169</sup>Tm (0.005), <sup>172</sup>Yb (0.02), <sup>175</sup>Lu (0.008), <sup>178</sup>Hf (0.014), <sup>181</sup>Ta (0.0035), <sup>208</sup>Pb (0.021), <sup>232</sup>Th (0.01), <sup>238</sup>U (0.005) at 20 ms integration time each.

## 3.4. XRF Bulk Rock Chemistry

An oven dry (105 °C) fused disc of each sample was measured by X-ray fluorescence spectrometry using a Robotic TGA system at the Bureau Veritas Mineral Laboratory in Perth (Australia) to detect the bulk rock elemental composition. Fourteen major elements and oxides (CaO, F, SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, Fe<sub>2</sub>O<sub>3</sub>, K<sub>2</sub>O, MgO, SO<sub>3</sub>, TiO<sub>2</sub>, P<sub>2</sub>O<sub>5</sub>, Na<sub>2</sub>O, MnO, Cl) given in wt.% were determined for low-sulphide samples and twelve major elements and oxides (Fe<sub>2</sub>O<sub>3</sub>, SiO<sub>2</sub>, Al<sub>2</sub>O<sub>3</sub>, TiO<sub>2</sub>, CaO, MnO, SO<sub>3</sub>, MgO, K<sub>2</sub>O, Pb, P<sub>2</sub>O<sub>5</sub>, ZnO) given in wt.% were determined for high-sulphide samples.

For error correction, internal rock standards of fluorspar, stream sediments, kinzingite and phosphate rock were used for the first determination. For the second determination, the internal rock standards OREAS 134b Zn-Pb-Ag, Gold-Enargite, Greenstone Ore, Canmet SU-1b and Canmet CPB-2 were used.

# 4. Results

# 4.1. Petrography

Hydrothermal fluorite found in drill core and hand samples from the investigated areas (i.e., ABB, FCB, LSB, and Harz Mountains) displays several modes of occurrence (Figure 2). In the ABB, fluorite is an abundant mineral phase that occurs as small interstitial grains (<100  $\mu$ m), as vuggy space-fillings, in massive mineralization associated with hydrothermal veins and as coatings on tectonic planes (Figure 2A,F–I). Hyp-idiomorphic grains (1–100s micron range) within ±quartz ± calcite ± anhydrite veins (Figure 2G) are common. In the LSB, hydrothermal fluorite is often associated with sulphides such as sphalerite and pyrite and occurs as rims surrounding sphalerite (Figure 2B) or as massive (mm-cm-scale) fluorite mineralization (Figures 2J and 3E,F) that is often associated with quartz, calcite, anhydrite and in rare cases, barite. The latter gangue minerals are also commonly present together with fluorite in veins and on tectonic planes in the volcanic and sedimentary host rocks in the FCB (Figure 2C,D,K,L) and Harz Mountains (Figure 2E,M,N). Although fluorite is in most cases colourless or only displays a slight hue, some distinctly coloured varieties occur (Figure 2A–E). Finely dispersed fluorite was identified with short wavelength UV light (Figure 2B). Anhydrite is often coeval with fluorite but also occurs together with paragenetically later barite in a sample from ABB Well D (Figures 2I and 3D).

# 4.2. SEM (BSE, CL)

Back-scattered electron and CL reveal complex textures in fluorite and associated hydrothermal minerals such as calcite and quartz (Figure 3A–F) that cannot be observed with transmitted or reflected light microscopy. Cyclical growth features (i.e., cyclical banding) are apparent in fluorite, calcite and quartz (Figure 3F). Furthermore, fine, vein-like textures within the fluorite are visible in several samples (Figure 3B). Figure 3E shows a high magnification CL image of a complex patchwork or network of distinct fluorite phases that have different luminescent characteristics. Larger patches of two distinct phases (I-lighter grey and II-medium grey, corresponding to I and II in geochemistry chapter), that were also analysed using LA-ICP-MS, are surrounded by thin rims of a third phase (darkest grey) which lie below the spatial resolution of the LA-ICP-MS.

# 4.3. Geochemistry

The collected LA-ICP-MS data for fluorite from the investigated areas are summarized in Tables 2 and 3 as well as in the corresponding boxplots and spidergrams in Figures 4 and 5.

## 4.3.1. Minor and Trace Elements (Excluding REY)

Minor and trace element concentrations (Figure 4) are characteristically low in fluorite from all areas—median of 633 ppm—i.e., the sum of all measured median element concentrations (excluding the REY) ranges between a minimum of 221 ppm (ABB Well DII) and a maximum of 2048 ppm (ABB Well DI). The later value for ABB Well DI is likely affected by an exceedingly high value for K which is based on only one measurement and likely attributed to a mineral inclusion that was overlooked during LA-ICP-MS data screening.



Figure 2. (A): Pinkish fluorite from ABB Well E. (B): Polished mount with fluorite surrounding sphalerite in hydrothermal quartz-calcite-sulphide vein from the Main Dolomite (Staßfurt Carbonate, Ca2) carbonate, LSB Well F. The right half of the polished mount shows the fluorescence of the fluorite (photographed after UV light exposure). (C): Pale violet fluorite in quartz-calcite vein from FCB Sohle 3 (see also K). Dark angular crystals are large chalcopyrite grains. (D): Greenish to colourless fluorite in quartz-calcite vein from FCB Sohle 4. (see also L) (E): Lavender fluorite in quartz vein (Biwender Vein) from the Harz Mountains. Dark areas are siliciclastic sediment clasts. (see also M). (F): Fluorite (dark) associated with calcite and anhydrite in a hydrothermal vein from ABB WELL A. (crossed polars, transmitted light) (G): Large hypidiomorphic fluorite (dark grey) associated with calcite and anhydrite. Quartz (bottom left) often occurs as idiomorphic inclusions in fluorite. (reflected light). (H): Massive fluorite mineralization with large clusters of hydrothermal anhydrite from ABB Well C. (transmitted light, crossed and plain polars). (I): Massive fluorite mineralization with late barite-anhydrite vein and small quartz veinlets from ABB Well D. (transmitted light, crossed and plain polars). (J): Fluorite associated with carbonate that displays recrystallizing textures is intergrown with organic material (dolomitic Ca2 carbonate) from LSB Well F. (transmitted light) (K): Quartz-calcite-fluorite vein in ignimbrite from FCB Sohle 3. (transmitted light, crossed polars). (L): Quartz-calcite-fluorite vein in ignimbrite from FCB Sohle 4. (transmitted light, crossed polars) (M): Quartz-fluorite vein (Biwender Vein) in fine-grained clastic sedimentary host rock. Quartz shows characteristic syntaxial candle growth texture. A set of later quartz veinlets is cross-cutting the quartz-fluorite vein. (transmitted light, crossed polars). (N): Hypidiomorphic fluorite (dark grey) in quartz-fluorite vein (Biwender Vein) from the Harz Mountains. (reflected light)-qz: quartz, fl: fluorite, cal: calcite, cb: carbonate mineral, anh: anhydrite, ba: barite, sp: sphalerite, sed: clastic sediment clast.



**Figure 3.** (**A**). BSE image of hydrothermal fluorite, quartz and anhydrite from ABB Well B. (**B**): Colour-CL image of hydrothermal fluorite, quartz and anhydrite. Sector zoning (generation\* I and II) is a prominent feature in fluorite. Anhydrite shows characteristic deformation lamellae. Quartz also displays growth textures that indicate a successive growth under varying conditions. (**C**): BSE image of a large fluorite grain with pronounced sector zoning and cyclical growth patterns. At least two distinct fluorite generations (I and II) can be observed in this sample from ABB Well C. (**D**): BSE image of a barite-anhydrite vein in fluorite from ABB Well D. Barite and anhydrite represent the last stage of this hydrothermal event. Due to the large contrast difference between barite and fluorite the growth patterns in fluorite are barely visible in this image. (**E**): CL image of fluorite from LSB Well A displaying the intricate network of fluid pathways and growth textures typical of complex sectoral growth (e.g., fir tree structures, see Discussion, Figure 7G). Two main generations (I and II) can be observed in this sample. \* 'generation of sectoral and oscillatory zoning (CL image, see Discussion, Figure 7F). Two distinct fluorite generations (I and II) can be observed in this sample. \* 'generation' is a term that in these cases does not necessarily denote an attribution to different fluid generations and a difference in age (as discussed in Section 5).

**Table 2.** LA-ICP-MS minor and trace element data (excluding REY) for fluorite (polished petrographic mounts) from the investigated areas, including total number of in-situ analyses per sample (#) and minimum (Min), maximum (Max) and median values for the sample populations. Null represents the number of analyses that returned values below the detection limit. All concentrations in parts per million (ppm).

ID and (#)	Stats	Li	Na	Mg	К	Mn	Fe	Zn	Ga	Sr	Ba	Pb
	Null	7	5	6	0	6	0	6	0	0	4	1
	Min	1.14	6.4	2.56	13.1	0.45	577	0.79	0.46	3.85	0.28	0.041
ADD Well AI (9)	Max	1.25	183	3.69	53.0	3.19	919	1	1.56	18.4	0.75	0.337
	Median	1.19	77.8	2.67	20.2	1.67	707	0.79	1.4	12.0	0.39	0.177
	Null	3	1	2	0	2	0	4	3	0	2	3
ARR Mall AII (5)	Min	1.97	17.7	2.59	14.8	0.99	693	5.13	0.051	7.29	1.5	0.385
Abb Well All (5)	Max	13.1	1583	106	327	25.7	907	5.13	0.114	56.6	6.1	2.66
	Median	7.56	140	7.32	28.4	3.86	741	5.13	0.082	11.17	1.53	1.52
	Null	8	6	14	2	7	0	9	16	0	5	16
A BB Moll BI (16)	Min	0.47	21.7	16	17.1	2.06	117	2.76	_	9.83	0.66	—
Abb well bl (16)	Max	48.3	3424	16.2	698	87.9	281	59.1	_	140	28	—
	Median	4.03	253	16.1	38.6	4.87	210	6.89	—	17.7	2.07	_
	Null	5	5	8	0	5	0	4	2	0	4	12
A DD $M_{2}$    DII (12)	Min	0.333	48	7.31	14.7	1.38	131	1.39	0.145	7.9	1.27	—
ADD Well DII (12)	Max	12	1153	14.4	271	27.5	206	22.8	0.925	77.1	9.37	_
	Median	2.65	212	10.9	28.7	7.67	164	3.91	0.392	13.8	2.99	_
	Null	5	3	20	1	5	0	10	20	0	5	11
A BR Moll CI (21)	Min	0.60	52.6	4.14	13.7	0.69	123	1.21	0.207	5.79	0.708	0.41
ABB Well CI (21)	Max	11.9	1045	4.14	158	19.4	323	4.74	0.207	1676	8.43	57.4
	Median	3.33	192	4.14	42.55	4.70	203	1.44	0.207	15.6	2.22	1.26
	Null	5	4	19	0	6	0	12	1	0	5	14
A PP Mall CII (24)	Min	0.403	31.1	5.71	15.7	1.19	160	1.21	0.16	4.53	0.677	1.01
Abb wen Cli (24)	Max	29	2142	9.58	357	35.7	343	7.5	2.47	77.9	15	44.9
	Median	3.52	310	7.49	58.7	6.33	219.5	2.04	1.07	17.9	2.38	3.83
	Null	1	1	1	0	0	0	1	1	0	0	1
	Min	_	_	_	1743	6.33	225	_	—	18.5	55.4	—
ADD Well DI (1)	Max	_	_	_	1743	6.33	225	_	—	18.5	55.4	—
	Median	_	—	—	1743	6.33	225	—	—	18.5	55.4	

Table 2. Cont.

ID and (#)	Stats	Li	Na	Mg	К	Mn	Fe	Zn	Ga	Sr	Ba	Pb
-	Null	3	3	3	0	2	0	3	1	0	1	3
	Min	_	_	_	22.6	30.5	62.3	_	0.378	12.7	1.22	_
Abb Well DII (3)	Max	—	—	—	61	30.5	256	_	0.854	15.5	2.03	_
	Median	—	—	—	28.9	30.5	146	—	0.616	12.9	1.625	—
	Null	1	0	1	0	1	0	2	3	0	0	0
A PR $M_{\rm oll}$ EI (4)	Min	0.214	6.5	1.91	15.17	0.322	647	1	0.03	26.6	0.211	0.152
ADD Well EI (4)	Max	1.24	258	776	49.2	245	729	7.55	0.03	5134	1.69	4.81
	Median	0.271	25.9	235	18.1	3.92	657	4.28	0.03	504	1.13	0.249
	Null	7	0	0	0	2	0	0	9	0	0	0
	Min	0.55	169	19.1	15.1	0.19	130	0.8	0.028	47.1	0.325	0.85
LSD Well FI	Max	10.5	9032	292	155	4.18	188	34.7	0.348	465	14.9	95.2
	Median	1.43	469	37.3	36.5	0.562	153	6.53	0.083	93.4	1.01	12.75
	Null	13	0	0	0	3	0	0	5	0	1	0
	Min	0.43	37.9	8.2	8.22	0.304	117	5.17	0.04	10.1	0.07	0.26
LSB well GI	Max	2	696	2170	285	106	182	2935	0.602	12000	3.49	63.4
	Median	0.89	216	29.2	37.3	0.95	154	90.8	0.223	31.2	0.77	14.0
	Null	6	0	0	0	2	0	0	9	0	0	0
Soble 3 Electringen High	Min	0.44	30.6	16	8.77	0.44	142	2.25	0.022	9.15	0.249	0.436
Some 5, Flechtingen Flight	Max	4.78	708	43.8	267	9.43	324	57.3	0.022	34.3	3.82	21.1
	Median	2.63	139	22.9	22.3	1.07	165	5.99	0.022	11.7	0.554	12.1
	Null	7	0	0	0	3	0	0	1	0	0	0
Soble 4 Electringen High	Min	1.04	52.3	8.53	10.4	0.313	168	0.83	0.039	8.78	0.117	3.13
Johne 4, Fleentingen Flight	Max	2.88	569	161	52.3	4.35	200	12.11	0.296	24.4	1.65	43.6
	Median	1.96	164	28.3	22.7	1.275	183	2.53	0.112	10.8	0.53	13.6
	Null	0	0	1	0	0	0	4	0	0	0	0
Piruan dan Vain I. Hang	Min	0.444	74.2	1.8	30.9	1.74	841	1.03	0.066	24.1	0.8	0.895
biwender veni i, narz	Max	4.56	728	20.2	186	15.2	1184	8.28	0.226	64.5	15.9	9.59
	Median	1.94	265	4.17	60.7	4.49	978	1.77	0.152	34.9	2.03	2.3
	Null	3	0	1	0	1	0	3	2	0	1	0
Rivender Vein II. Herr	Min	0.366	5.92	1.57	18.0	0.55	884	0.85	0.031	10.8	0.197	0.389
biwender vent if, Harz	Max	0.366	37.4	2.91	32.8	0.98	1135	0.85	0.059	19.9	3.37	5.51
	Median	0.366	20.1	2.21	27.1	0.564	984	0.85	0.045	12.8	0.94	3.03

**Table 3.** REY concentrations in fluorite (polished petrographic mounts) from the investigated areas, including number of in-situ analyses per sample (#), minimum (Min), maximum (Max) and median values. Null represents the number of analyses that returned values below the detection limit. All concentrations in parts per million.

ID and (#)	Stats	La	Ce	Pr	Nd	Sm	Eu	Gd	Тb	Dy	Ŷ	Но	Er	Tm	Yb	Lu
	Null	0	0	0	0	0	0	0	0	0	0	1	0	2	6	7
$\Delta BB W_{0} = 1 \Delta I (9)$	Min	17.3	22.27	2.17	9.59	0.646	0.024	0.295	0.014	0.069	1.17	0.0255	0.031	0.005	0.041	0.015
$\operatorname{Hbb}\operatorname{Hch}(\mathcal{I})$	Max	55.4	96.96	9.43	32.2	4.5	0.459	2.56	0.3	1.68	35.9	0.334	0.655	0.049	0.14	0.0212
	Median	40.1	71.46	6.51	25.2	2.88	0.354	1.88	0.153	0.91	16.8	0.132	0.261	0.0262	0.104	0.0181
	Null	0	0	0	0	0	0	0	0	0	0	0	0	1	1	3
ABB Well AII (5)	Min	0.113	0.259	0.049	0.4	0.413	0.09	0.85	0.188	1.62	53.8	0.399	0.873	0.0533	0.232	0.027
	Max	1.38	2.78	0.307	1.85	1.96	0.486	4.67	0.97	7.69	119	1.46	2.45	0.229	0.72	0.059
	Median	0.568	1.43	0.211	1.36	0.77	0.374	2.72	0.542	4.21	80.8	0.912	1.61	0.178	0.539	0.043
	Null	7	8	12	13	13	13	10	8	2	0	4	8	14	16	16
ABB Well BI (16)	Min	0.082	0.076	0.046	0.216	0.298	0.06	0.577	0.063	0.296	12.6	0.059	0.172	0.049	—	—
1122 (10)	Max	0.328	0.382	0.1	0.624	1.13	0.215	1.63	0.292	1.58	65.3	0.326	0.688	0.06	—	
	Median	0.206	0.182	0.0555	0.498	0.556	0.191	1.055	0.185	0.533	22.85	0.17	0.392	0.0545		
	Null	1	1	1	1	3	2	4	6	5	0	8	9	12	12	12
ABB Well BII (12)	Min	3.39	3.94	0.166	1.32	0.294	0.044	0.365	0.052	0.163	0.58	0.047	0.125	_	—	_
()	Max	26.2	38.2	4.26	21.3	2.79	0.331	2.58	0.121	0.61	29.8	0.117	0.349		—	
	Median	13	18	2.36	9.73	1.46	0.158	0.876	0.0955	0.201	8.05	0.0505	0.172			
	Null	3	1	5	5	7	6	4	4	2	0	3	4	5	6	13
ABB Well CI (21)	Min	0.139	0.071	0.035	0.238	0.262	0.075	0.337	0.075	0.303	2.36	0.059	0.439	0.04	0.293	0.033
	Max	2.5	4.24	0.805	4.3	3.09	0.75	7.15	1.82	12.1	152	1.96	4.45	0.266	0.919	0.108
	Median	0.544	0.908	0.164	1.005	0.916	0.304	2.67	0.644	4.36	88	0.843	1.92	0.188	0.487	0.051
	Null	0	0	0	0	2	4	3	4	3	0	4	5	14	15	22
ABB Well CII (24)	Min	4.74	5.17	0.771	2.69	0.328	0.079	0.403	0.072	0.199	0.679	0.051	0.111	0.039	0.14	0.03
	Max	81.4	102	14.4	49.3	7.57	1.06	5.39	1.06	6.11	120	1.23	2.34	0.179	0.54	0.035
	Median	38.6	48.1	6.72	21.05	2.59	0.428	2.26	0.283	1.57	27.75	0.275	0.602	0.091	0.34	0.0325
	Null	0	0	1	0	1	0	1	1	0	0	1	1	1	1	1
ABB Well DI (1)	Min	0.954	1.03	—	0.971	—	0.241	_	_	0.908	36.6	_	—	—	—	—
TIDD Well DI (1)	Max	0.954	1.03	—	0.971	—	0.241	—	—	0.908	36.6	—	—	—	—	—
	Median	0.954	1.03		0.971		0.241			0.908	36.6		_		_	
	Null	0	0	0	0	1	2	2	2	2	0	2	2	3	2	3
ABB Well DII (3)	Min	8.56	9.53	1.46	6.67	0.768	0.256	1.64	0.246	1.57	4.28	0.405	1.08	—	1.28	—
1100 Well Dil (3)	Max	17.4	29.9	3.16	8.98	1.59	0.256	1.64	0.246	1.57	69.8	0.405	1.08	_	1.28	_
	Median	8.59	16.2	2.24	7.33	1.179	0.256	1.64	0.246	1.57	7.83	0.405	1.08		1.28	—

Table 3. Cont.

ID and (#)	Stats	La	Ce	Pr	Nd	Sm	Eu	Gd	Tb	Dy	Ŷ	Ho	Er	Tm	Yb	Lu
	Null	0	0	0	0	1	3	0	1	1	0	1	2	2	1	3
	Min	0.018	0.0338	0.0137	0.054	0.104	0.0105	0.044	0.0114	0.085	0.029	0.0109	0.025	0.0054	0.018	0.0114
ADD Well EI (4)	Max	0.287	0.565	0.128	0.685	0.193	0.0105	0.159	0.0503	0.201	1.068	0.0348	0.094	0.0076	0.14	0.0114
	Median	0.0539	0.0767	0.0607	0.353	0.109	0.0105	0.15	0.021	0.163	0.4025	0.0129	0.0595	0.0065	0.035	0.0114
	Null	0	0	0	0	2	0	1	2	0	0	3	2	6	9	9
I SB Wall EI (16)	Min	0.0237	0.087	0.0167	0.088	0.026	0.0072	0.036	0.0061	0.032	0.678	0.0077	0.013	9.60E-04	0.0125	0.0008
LOD WEIT I'I (10)	Max	0.081	0.213	0.0386	0.254	0.1	0.0282	0.112	0.018	0.099	1.544	0.02	0.044	0.013	0.035	0.0105
	Median	0.0514	0.151	0.0272	0.163	0.045	0.014	0.071	0.0123	0.0625	1.1125	0.011	0.025	0.00515	0.021	0.0037
	Null	4	1	6	12	13	12	10	10	14	1	11	9	10	9	9
I CD Wall CI (16)	Min	0.0008	0.0047	0.0048	0.0038	0.0083	0.0042	0.014	0.002	0.024	0.01	0.0012	0.0036	0.0018	0.0055	0.002
LSD Well GI (16)	Max	0.501	0.956	0.107	0.643	0.065	0.018	0.087	0.014	0.027	0.461	0.006	0.031	0.0083	0.092	0.0062
	Median	0.0356	0.0213	0.0118	0.045	0.017	0.0118	0.019	0.00245	0.0255	0.095	0.0037	0.0083	0.00485	0.0108	0.0045
	Null	1	0	1	0	1	0	0	0	0	0	0	0	1	0	4
Sohle 3, Flechtingen	Min	0.0045	0.0172	0.0035	0.01	0.035	0.0325	0.308	0.049	0.496	32.7	0.078	0.211	0.0193	0.023	0.0135
High (10)	Max	0.239	0.462	0.0606	0.451	0.502	0.384	1.87	0.458	3.12	138	0.555	1.43	0.097	0.383	0.0377
-	Median	0.0493	0.0598	0.0153	0.0575	0.095	0.0915	0.789	0.164	1.30	75.6	0.303	0.701	0.0494	0.19	0.0226
	Null	0	0	0	0	0	1	0	3	3	0	2	1	5	5	4
Sohle 4, Flechtingen	Min	1.53	2.321	0.236	1.037	0.061	0.0148	0.024	0.0064	0.031	0.725	0.0052	0.0076	0.00094	0.014	0.00094
High (9)	Max	9.46	18.7	2.2	9.39	1.43	0.159	0.45	0.0159	0.112	4.51	0.0249	0.039	0.0045	0.021	0.0069
-	Median	4.89	7.78	0.775	3.09	0.32	0.0444	0.114	0.012	0.047	1.342	0.0088	0.0188	0.00375	0.0175	0.0031
	Null	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0
Biwender Vein I,	Min	0.955	2.84	0.468	3.11	1.63	1.01	3.55	0.621	4.15	83.9	0.919	1.94	0.226	1.21	0.08
Harz (9)	Max	1.94	6.03	1.11	7.45	5.16	2.78	12.02	2.401	15.87	188.8	2.9	6.71	0.702	3.35	0.297
	Median	1.26	4.38	0.715	5.82	3.65	1.69	8.64	1.612	10.12	128	2.064	4.47	0.463	2.14	0.237
	Null	0	0	0	0	1	1	0	1	0	0	0	0	0	0	1
Biwender Vein II,	Min	0.084	0.14	0.0277	0.152	0.177	0.204	0.155	0.18	0.329	9.95	0.037	0.104	0.023	0.082	0.0096
Harz (4)	Max	0.215	0.423	0.08	0.783	0.618	0.389	1.54	0.216	1.56	23.7	0.304	0.459	0.05	0.211	0.0394
	Median	0.168	0.234	0.0557	0.413	0.455	0.252	0.9	0.209	1.345	19.7	0.236	0.415	0.0454	0.151	0.0204



**Figure 4.** Minor and trace element concentrations (excluding REY) in fluorite from the different investigated localities. Numbers in the boxplots indicate the number of analyses (above the level of detection) the corresponding values are summarized in Table 2. Note that the displayed median and mean values are based on the logged data and may therefore deviate from the data in Table 2.



**Figure 5.** C1 chondrite-normalized, after [76] REY<sub>CN</sub> plots for fluorite (**a**–**g**) and whole rock samples (**h**,**i**) from ABB (**a**–**d**), LSB (**e**), Flechtingen High, FCB (**f**), and Biwender Vein (Harz) (**g**). Median values are represented by the dashed lines. Compositional ranges for the Biwender Vein and the ABB and LSB whole rock data are plotted as references in the background in plots where applicable. Colour coding corresponds to Figure 4. The corresponding REY concentrations are listed in Table 3.

Only Na and Fe concentrations reach median concentrations that consistently exceed 100 ppm across all sample sites. Only samples ABB Well AI, AII, EI and BVII (Biwender Vein II) have median concentrations below 100 ppm for Na. Lithium contents are comparable among all sample groups and median concentrations range between 0.3 and 7.5 ppm. Sodium and potassium concentrations are also relatively evenly distributed and show comparable ranges in most samples-median values of 140–470 ppm and 20–60 ppm, respectively. Manganese shows slightly higher median concentrations in samples from the ABB (1.67–30.5 ppm) than in the other sample populations (0.5–1.3 ppm). Only BVI (Biwender Vein I) has a comparable median Mn concentration (4.49 ppm). Lead shows the opposite trend with median concentrations being about an order of magnitude higher in samples from the LSB and Flechtingen High compared to the ABB and BV fluorites. Iron shows the most pronounced and characteristic variation among the investigated sample groups. Median concentrations of 670-984 ppm can be found in samples ABB Well AI, II and CII as well as in the two sample populations from the BV. The remaining samples have significantly lower median concentrations of between 150–220 ppm. Zinc, Ga and Ba show no distinct compositional trends among the different fluorite populations. Gallium concentrations are commonly below the level of detection or slightly above with median concentrations barely exceeding 1 ppm in only a couple of samples. The high Ba concentration for sample ABB well DI is likely attributed to a mineral inclusion (see also K content). Strontium concentrations plot consistently at median values of around 10–20 ppm. Samples ABB Well EI, LSB Well FI, GI and BVI show greater ranges and or higher median concentrations of up to ca. 500 ppm.

There appears to be a subtle but noticeable trend between the I and II fluorites from ABB Well A and C, with generation II displaying slightly higher median minor and trace element contents. However, this difference is always less than an order of magnitude. Fluorite from the Biwender Vein displays the opposite trend with higher concentrations of all minor and trace elements except Fe and Pb in BV I compared to BVII. This trend is also reflected in the REY concentrations (Figure 5g).

#### 4.3.2. Rare Earth Elements and Yttrium (REY)

Similar to the overall minor and trace element contents the REY concentrations are low in comparison to fluorite from magmatic-hydrothermal settings. The median sum of REY ranges from 0.3 to 176 ppm. Despite these comparatively low concentrations, characteristic trends and patterns are evident in the REY signatures of fluorite from the different sample sites (Figure 5). While fluorite from the LSB shows a flat REY<sub>CN</sub> pattern (Figure 5d), fluorites from the ABB show two distinct REY<sub>CN</sub> patterns (Figure 5a–d). The two generations are labelled I and II in the corresponding spidergrams in Figure 5. Generation I is consistently enriched in LREE (La to Sm) and shows a significant depletion in the HREE compared to fluorite generation II (Figure 5a–d). These trends are particularly pronounced in fluorite from ABB Well A, B and C. Fluorite from Sohle 3 and 4 (Flechtingen High, FCB) show an almost identical signature with Sohle 4 corresponding to ABB generation I and Sohle 3 corresponding to ABB generation II. Due to a limited number of analyses the trend of LREE enrichment and HREE depletion in chondrite-normalized patterns is less pronounced but still visible in fluorite from ABB Well D (Figure 5d). Fluorite from ABB Well E displays a flat C1 chondrite-normalized signature and is overall quite similar to the signatures found in fluorite from the LSB (Figure 5d,e) which also correspond with previously published data by Duschl et al. [12]. Negative Eu<sub>CN</sub> anomalies were also in some of the samples (ABB Well AI, CI, EI) but are not particularly pronounced. A pronounced positive Y<sub>CN</sub> anomaly is present in most samples. However, fluorite from ABB Well DI, EI and LSB Well GI does not show this relative Y enrichment in the median concentrations. Fluorite from the Biwender Vein can be divided into two groups. Both show a similar shape, with a noticeable relative depletion in LREE (La-Nd) and HREE (Er-Lu), but differ in their relative REY concentrations, that is, BVI largely plotting above 10 and BVII plotting below 10 in the C1 chondrite-normalized spidergram (Figure 5g).

The whole rock (i.e., bulk drill core) REY data from the ABB and the LSB are shown for comparison in Figure 5. Both display a minor relative enrichment in LREE in comparison to HREE with the overall

REY content in the ABB whole rock data being an order of magnitude higher than those detected in the LSB whole rock samples. A negative  $Eu_{CN}$  and a positive  $Y_{CN}$  anomaly is discernible in the ABB whole rock data. Note that the ABB REY whole rock range is very similar in overall ratio and shape to ABB fluorite type I.

# 5. Discussion

#### 5.1. Fluorite Precipitation and the Fluorine Source

Fluorite in the deeply covered Paleozoic units of the NGB can be of sedimentary or hydrothermal origin. A syn-sedimentary formation of fluorite in the NGB has been ascribed by Möller et al. [25]. They also discussed the processes leading to the necessary enrichment of fluorine from marine waters to form fluorite. Seawater and marine pore waters are usually under saturated in F- and thus precipitation of fluorite is prevented in oxygenated seawater. Under these conditions the diagenetic formation of fluorite is linked to dolomite formation in calcareous sediments due to Mg<sup>2+</sup> removal from pore fluids and consequent release of F<sup>-</sup> due to break-up of MgF<sup>+</sup> complexes [25]. Organic material e.g., [77,78], fresh water influx with elevated fluorine concentrations attributed to pegmatitic sources [79], or a volcanogenic source [80,81] have also been invoked to account for the required saturation to allow fluorite precipitation under diagenetic conditions. It is also possible that diagenetic fluorite may act as a precursor for later hydrothermal fluorite generations, that is, a remobilization of primary sedimentary fluorite yielded sufficient fluorine for the deposition of hydrothermal fluorite. Basinal brines that migrated through igneous units (i.e., thick successions of Rotliegend volcanics in the ABB and FCB) may also have acquired the necessary fluorine contents to achieve hydrothermal saturation and deposition along structurally controlled veins within the NGB. In the LSB zones of hydrothermal sulphide mineralization, specifically sphalerite and galena, have been documented in Zechstein carbonates by several authors i.e., [2,82-84]. Although researchers have argued about the possible sources of the mineralizing fluids and the style of mineralization (from magmatic to MVT) [82,85–92], recent studies have presented convincing evidence that structurally and thermochemical sulphate reduction (TSR)-controlled hydrothermal basinal brines led to the Pb-Zn mineralization and the associated fluorite-barite mineralization [2]—a model that can be equally applied to the hydrothermal fluorite mineralization in the ABB, although a different lithology (i.e., thick volcanic layers) translates into distinct variations compared to the LSB [14,93].

#### 5.2. REY Composition and Systematics

In general, fluorites preserve or inherit the REY signature of the hydrothermal fluid from which they precipitated and hence offer an insight into the source of the REY in the hydrothermal fluid (i.e., intrusive or volcanic rocks) e.g., [36]). In particular in the ABB, where a suite of volcanogenic units of variable compositions [7] is part of the basin's Paleozoic rock inventory, the possibility that percolating basinal brines migrated through these volcanic rocks along steep and deep-reaching faults [94] and picked up their distinct REY signature, including a typical negative Eu anomaly, is a plausible explanation for the observed REY<sub>CN</sub> signatures. Lüders et al. [12], for example, provide evidence from fluid inclusion investigations that indicated high water-rock interactions with Permian volcanic and sedimentary rocks in the ABB. However, a primary volcanogenic fluid source can be discarded for both the ABB and LSB [2,12,18,93]. The flat REY<sub>CN</sub> signature [95,96] and the overall low concentrations are strong indicators that basinal brines derived from marine water represent the mineralizing fluid in the LSB. A comparison of a selection of whole rock volcanic REY<sub>CN</sub> patterns with median ABB fluorite signatures is shown in Figure 6. The median fluorite ABB I signature displays a similar REY<sub>CN</sub> pattern to the whole rock volcanic rocks from the NEGB compiled by Benek et al. [7], with a relative enrichment of LREE. Particularly the rhyolite is a good match for the LREE to MREE. However, while the volcanics show a flat HREE pattern the fluorite ABB I generation displays a slight slope downwards toward the heavier HREE and a distinct positive peak for Y, which is missing

entirely in the volcanics. Fluorite ABB II bears no similarities with the volcanic signatures, that is, is significantly depleted in LREE in comparison.



**Figure 6.** Comparison of C1 chondrite-normalized, [after 76] REY<sub>CN</sub> data for volcanic whole rock samples from the NEGB, [after 7] and the median fluorite signatures for generation I (AI, BI, CI, DI) and II (AII, BII, CII, DII) from the ABB (see also Figure 5).

Many authors have suggested that fluoro-complexes as the primary means of REE transport and the deposition of fluorite [97,98]. However, Migdisov and William-Jones [99] presented experimental evidence that challenges this assumption. They argued that fluoro-complexes are in fact negligible for the hydrothermal transport of REE and that chloride- and sulphate-complexes are much more efficient at binding and transporting REE in hydrothermal fluorine-bearing fluids whereas fluoro-, carbonate-, and phosphate-complexes are important as depositional ligands [35]. Lee and Byrne [100,101] presented stability constants for chloro-complexes which exhibit a decrease from LREE to HREE in accordance with the trends observed in our data. Furthermore, Möller et al. [33] noted that LREE are more easily incorporated into fluorite due to their similar radii compared with Ca<sup>2+</sup>, leading to a preferential incorporation and a positive LREE anomaly in many early generation fluorites. Later generations consequently are relatively depleted in LREE if precipitated by the same fluid [33], especially if the REE are not bound in fluoro-complexes [102]. Following Möller's interpretation, the fluorite populations in this study that show relatively lower LREE concentrations would therefore have to be considered as a later or secondary (remobilized) fluorite generation (II) compared to the fluorite (generation I) with relatively elevated LREE contents. However, adsorption, chemical complexation, changes in temperature and salinity can lead to highly localized variations in fluorite compositions including the REE [30,31,36,38], making it difficult to attribute different REE signatures to single controls. Moreover, our petrographic observations and characteristic growth patterns detailed in Sections 4.2 and 5.3 strongly suggest that both fluorite generations formed during one hydrothermal event and reflect micro-scale adsorption features and thermodynamic and compositional variations of the precipitating fluid. Schwinn and Markl [36] proposed that relative LREE enrichment in the outer zones of fluorite crystals may represent a sub-micron LREE-enriched phase or can be attributed to LREE clustering in later, cooler fluids. However, adsorption favours HREE, whereas LREE are more easily held in solution by chemical complexation [32,100,101]. Diffusive processes are highly unlikely to have contributed to compositional variations—as primary sector zoning and oscillatory zonings are preserved [37]—especially at the low formation temperatures (<~250 °C) observed in the ABB and LSB.

Several trends are evident in the chondrite-normalized REY<sub>CN</sub> spidergrams in Figure 5. The elevated REY concentrations observed in fluorites from the ABB, the FCB and the Harz mountains provide evidence that they were formed from fluids that differed significantly from those in the LSB. Variations in complexation behaviour/chemistry and fluid-rock interaction may also have led to the distinctly different REY<sub>CN</sub> patterns, that is, a carbonate buffered system in the Permian shelf-carbonate dominated LSB versus Paleozoic volcanics and siliciclastic sedimentary rocks in the ABB, FCB and Harz. The minor but noticeable  $Eu_{CN}$  anomalies in our samples could be due to (a) an inherited signature that reflects the source REE (i.e., Rotliegend volcanics, Figure 6) e.g., [98], (b) adsorption features and crystallographic controls (see Section 5.3), (c) strongly reducing and or oxidizing conditions, or temperatures above 200 °C—thermochemical reduction of Eu<sup>3+</sup> to incompatible Eu<sup>2+</sup> [30,36,38], (d) analytical fractionation or bias due to the poor ablation behaviour of fluorite, or (e) a feature induced by the C1 chondrite). Option (c) is unlikely because there is no evidence for these conditions in the literature or in our investigation. Option (d) can be largely ruled-out because that fractionation should be consistent and affect all analyses, which is not the case.

The fluorite samples from the ABB and the Flechtingen High show a pronounced compositional variation between generation I and II, which corresponds to petrographic observations in the corresponding thin sections and polished mounts. Fluorite generation I (light under SEM) exhibits a trend that matches the whole rock composition (ABB whole rock REE). Fluorite generation II (dark under SEM) has distinctly lower LREE concentrations and aligns in terms of the overall trend with the hydrothermal vein fluorite from the Biwender Vein (Harz). These trends could be the result of a remobilization process accompanied by significant fluid/rock interaction with the surrounding sedimentary rocks or reflect physicochemical variations and intrinsic crystallographic controls within a single mineralizing event.

## 5.3. Remobilization versus Cyclical Growth or Sectoral Zoning

Cathodoluminescence and BSE images of hydrothermal fluorites (Figure 3) from the investigated areas reveal several prominent zoning features visualized as colour contrasts. Different types of growth zoning, mainly concentric and sectoral zoning, are illustrated and discussed further in Figure 7. As demonstrated by Marshall [72], CL in hydrothermal fluorite is mostly activated by intrinsic effects, that is, the incorporation of impurity atoms such as REE or trace elements such as Pb, Mn, Fe, Sr, and Y. Selective incorporation of foreign cations, above all REY, into the fluorite crystal lattice depending on crystal habits, and therefore effects of the surface free energy of specific crystal faces exposed to the hydrothermal fluid during precipitation, has been explained in detail by Bosze and Rakovan [40] and Rakovan [103]. These authors showed how this process is responsible for the fractionation of REE and trace elements during crystal growth, which may explain the significant differences in REE contents, especially LREE, found within individual fluorite crystals (i.e., generation I and II) of our samples. Type and quantity of incorporated cations that substitute for Ca<sup>2+</sup> in the crystal lattice depend on ionic charge and radius, element availability in the fluid, as well as the crystal morphology (e.g., octahedron, rhombic dodecahedron, cube etc.) (Figure 8). Crystal habits of hydrothermal fluorite and its variation during growth, however, may be a function of formation temperature and pH/Eh conditions [23,39,104]. Thus, the application of the Tb/Ca-Tb/La diagram [33] (Figure 9), which is widely used to distinguish between different parental fluids is problematic in cases where bulk analyses of fluorite are considered for genetic ore interpretation. Remobilization leads to an increasing Tb/La ratio by maintaining the same Tb/Ca ratio [29,33] and is a common feature in sedimentary and hydrothermal fluorite-bearing deposits e.g., [41]. However, this assumes a dominant complexation by F-ions which will keep HREE in solution during migration. The Tb/La versus Tb/Ca plot in Figure 9 illustrates that both fractionation during crystallization and remobilization trends are evident in the fluorite samples from the ABB, LSB, and Flechtingen High—if we consider the initial statement to be valid. For example, the distribution of ABB Well C (light) shows a spread parallel to the fractionation

trend proposed by Möller et al. [33]. Following this interpretation ABB Well A and C (light) and Flechtingen High Sohle 4 can be considered to represent the first fluorite stage. ABB Well A and C (dark) and Flechtingen High Sohle 3 appear to be a remobilization product of this first fluorite generation respectively.

The same trends can be observed in sample ABB Well B where a light and dark variety (SEM shades) was detected (Section 4.2). The LSB fluorites have low Tb/Ca ratios and moderate Tb/La ratios which places them in the sedimentary field. LSB Well A shows a data spread that is interpreted as a fractionation trend but none of the LSB fluorites seem to exhibit a remobilization trend that is evident in the Tb/La versus Tb/Ca plot. Both LSB fluorite samples plot close to fluorite measured by Duschl et al. [11] who also investigated fluorite in the LSB. Fluorite samples from the Biwender Vein (Harz) plot in the same area as the generation II ABB fluorite.



Figure 7. Various examples of growth zoning features typically found in hydrothermal fluorites as evident from the petrography and cathodoluminescence and BSE imaging (Figures 2 and 3). Different colours illustrate image contrasts, and thus, differences in chemical composition (corresponding to fluorite generation I and II). Plane section for (D,G,H) is |110|, all other examples are shown parallel to |001|. No apparent zoning (A) suggests crystallization under constant hydrothermal conditions, whereas concentric zoning (B) indicates discontinuous changes in the fluid chemistry over time; (C) is a typical representation of short-lived oscillating changes in the fluid chemistry. (D,E) Typical sectoral zoning in fluorite crystals with complex crystal habits ((D): cuboctahedral, (E): cubic + rhombic dodecahedral faces) that is only visible in specific plane sections. (F) Combination of sectoral and oscillatory zoning in a cubic fluorite crystal. (G) 'Fir-tree' zoning occurs when growth conditions (T/Eh) alternate over time and thereby favour different crystal morphologies (cuboctahedron  $\leftrightarrow$  cube), which proves pulsed crystallization (e.g., [106]). (H) 'Petal-type' zoning is the result of a continuous change from octahedral growth to cubic growth; the red line marks the beginning of shape transition (modified from [104]). Image (**B**,**E**) correspond to features visible in Figure 3B (parallel to |111|) and Figure 3C (parallel to |100|); image (F) resembles zoning details visible in Figure 3F, while image (G) is similar to growth patterns apparent in Figure 3E in the area labelled II.



Figure 8. (A) Schematic illustration of an idealised cube (grey) and octahedron (blue), respectively. Major crystallographic planes are indicated with respective indices, the section plane used for images in (B,C) is marked with green lines. (B) Assuming a fluid of constant composition, at higher temperatures fluorite will precipitate in octahedral shape (I) with only |111| planes exposed to the surrounding fluid (blue shade), which results in the preferential incorporation of elements of specific size/charge (compare [23]). Cooling of the fluid will now favour a cubic fluorite morphology and, consequently, the continuous formation of |100| planes (II), which show different sorption properties compared to 1111 planes (grey shade). Although precipitating synchronously from the same fluid, two different fluorite generations will form, since the two exposed crystal planes preferentially incorporate different elements. Note, how isochrones of fluorite growth (red lines) crosscut both generations; they represent former crystallographic growth surfaces that were exposed to the hydrothermal fluid at the same time. Continuous cooling will reduce the total area of |111| planes (II-III, cuboctahedron) and finally result in a purely cubic crystal shape (IV). In this example, temperature is the governing factor for crystal morphology and thus, for element incorporation. Thereby, it overrules other factors that usually control geochemical composition such as fluid composition itself, which may only play a minor role here (i.e., REE and trace element supply). (C) Repeated heating or influx of hot hydrothermal fluid may then cause a reversed growth cycle, when higher ambient temperatures eventually favour octahedral fluorite growth, resulting in the formation of |111| planes (V-VII). (D) Three-dimensional illustration of various fluorite morphologies during growth. Please note that zoning features observed in BSE or CL strongly depend on the section plane of the sample. Features presented in the above illustrations are only visible when the zoned crystal is observed normal to the section plane indicated in Figure 7A.

A remobilization or recrystallization trend is also apparent in the La/Ho versus Y/Ho plot in Figure 10. All fluorite samples show two distinct generations (I and II) that follow a remobilization trend [31]. Y/Ho ratios remain constant whereas La/Ho ratios decrease during remobilization.

Schwinn and Markl [36] pointed out that petrographically late fluorite from several of their investigated deposits display geochemical characteristics that reflect new fluid influx rather than remobilization. However, they also acknowledged that variable degrees of supersaturation and fluid/fluorite ratios as well as fluid flux lead to a complex precipitation process that reflects these

variable conditions. Hence, a rapid, pulsed crystallization on a micro-scale may display similar features to those seen when new (or later) fluids enter the system.



**Figure 9.** Tb/La versus Tb/Ca atomic ratio diagram after Möller et al. [33] for fluorite from the ABB, LSB, Biwender Vein (Harz), and Flechtingen High. For reference, fields compiled and established by Makin et al. [28], Duschl et al. [11], and Kraemer et al. [43]—North Pennine Ore Field (NPO), Alston Block (A) and Askrigg Block (B) are plotted. Labels, groupings and colours in the diagram (e.g., AI, Sohle 3) correspond to sample populations shown in Figures 4 and 5. See text for discussion.



**Figure 10.** La/Ho versus Y/Ho plot for a selection of fluorite samples from the ABB and the Flechtingen High. The grouping and colours correspond to those in Figure 5. See text for discussion.

Under certain conditions it is possible to produce two geochemically distinct types of fluorite from one and the same parental fluid within the same crystal (even without remobilization), in particular with regard to REE and trace elements. When sorption effects along crystal growth surfaces are responsible for impurity atom incorporation, crystal morphology during a specific growth event will control the type and amount of the respective impurity adsorption (Figures 7 and 8). As shown by Dill and Weber [23], among others, fluorite morphology locally strongly depends on fluid temperatures and hence fluid viscosity. For example, if |100| planes of cubic fluorites favour the incorporation of LREEs and yet cubic fluorite is more likely to form at relatively low temperatures, fluorite precipitation at low ambient temperatures will result in fluorite with low LREE concentration. Conversely, higher temperatures will produce octahedral fluorite crystals with comparatively higher LREE concentrations, regardless of the REE content provided by the parental hydrothermal fluid (as long as REEs are present). Since transformation from octahedral to cubic fluorite is a continuous process both fluorite types can coexist (see Figures 3 and 7). Fluctuations in temperature may then cause complex fir tree-type patterns as shown in Figure 7G.

In our samples, a possible third generation of fluorite is evident in the CL images (Figure 3E) as a dark, thin rim surrounding fluorite generation I and II along grain boundaries. The dark CL colour is thought to be attributed to high Sr contents which are concentrated in the residual fluid until it falls out as the last phase. At this stage, we do not have the analytical evidence to back this claim because this third generation eludes the spatial resolution of the applied analytical techniques.

# 6. Conclusions

Fluorites from the different investigated areas show characteristic petrographic features such as zoned growth, cyclical zoning and other complex zoning structures (i.e., fir tree structures). Petrographically discernible zones can also be discriminated geochemically and show signatures that suggest two (or more) distinct fluorite generations. Hereby, fluorite from the ABB and the FCB display REY<sub>CN</sub> patterns that invoke a similar fluid source and formation conditions (i.e., basinal brines that interacted with Lower Permian volcanic rocks at similar temperatures), whereas fluorite

from the LSB display flat REY<sub>CN</sub> signatures, comparable to the LSB whole rock patterns, that point to basinal brines derived from marine waters (pore waters) with no volcanic component (primary magmatic or through fluid-rock interaction) as the mineralizing fluids. A primary hydrothermal mineralization and a secondary remobilization geochemical signature is apparent in the frequently used Tb/La versus Tb/Ca and La/Ho versus Y/Ho discrimination diagrams for fluorite from the ABB and FCB. Our investigation suggests that the same trends are the result of zoned growth during a single mineralizing event rather than representing remobilization (i.e., secondary processes). This has implications for the interpretation of such trends and hence the inferred genetic evolution of fluorite that displays such geochemical patterns. The complex micro-scale intergrowth of these generations stresses the need for detailed petrographic investigations when geochemical data are collected and interpreted.

Author Contributions: For research articles with several authors, a short paragraph specifying their individual contributions must be provided. The following statements should be used "Conceptualization, P.N. and R.K.; Methodology, P.N., F.D. and M.R.; Validation, P.N., M.R. and F.D.; Formal Analysis, P.N., F.D. and M.R.; Investigation, P.N., M.R., F.D. and D.K.; Resources, R.K. and P.N.; Data Curation, P.N., M.R., F.D., D.K. and M.S.; Writing-Original Draft Preparation, P.N., M.R., F.D., D.K. and M.S.; Writing-Review & Editing, P.N., M.R., F.D., R.K., D.K. and M.S.; Supervision, P.N. and R.K.; Project Administration, P.N. and R.K.; Funding Acquisition, R.K."

Funding: This research was funded by the German Ministry for Eductaion and Research (BMBF), grant number [033R165].

**Acknowledgments:** We thank Exxon Mobile, Engie and Wintershall for access to their drill cores, and Norddeutsche Naturstein GmbH for access to the Flechtingen quarry. The authors also thank Helene Brätz for in-house LA-ICP-MS measurements and data integration. Three anonymous reviewers helped to improve the original manuscript greatly and we very much appreciate their comments and suggestions.

**Conflicts of Interest:** The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

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